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| | - | • | | | | |
| | (54) Title: BORONIC ACID AND ESTER INHIBITORS | OF TH | ROMBIN | | | |
| (57) Abstract | | | | | | |
| Novel boronic acid derivatives of formula (I), which are useful inhibitors of trypsin-like enzymes, are disclosed: R¹-Z-CH | | | | | | |
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<u>Title</u>

Boronic Acid and Ester Inhibitors of Thrombin

Field of the Invention

5 This invention relates to the discovery of new boronic acid derivatives which are inhibitors of thrombin and pharmaceutical compositions thereof.

Background of the Invention

- Hemostasis is the normal physiological process in which bleeding from an injured blood vessel is arrested. It is a dynamic and complex process in which thrombin plays a key role. Blood coagulation may occur through either of two cascades of zymogen activations, the
- extrinsic and intrinsic pathways of the coagulation cascade. The last protease in each pathway is thrombin, which acts to hydrolyze four small peptides (two FpA and two FpB) from each molecule of fibrinogen, thus deprotecting its polymerization sites. Once formed, the
- linear fibrin polymers may be cross-linked by factor XIIIa, which is itself activated by thrombin. In addition, thrombin is a potent activator of platelets, upon which it acts at specific receptors. Thrombin activation of platelets leads to aggregation of the
- cells and secretion of additional factors that further accelerate the creation of a hemostatic plug. Thrombin also potentiates its own production by the activation of factors V and VIII (see Hemker and Beguin in: Jolles, et. al., "Biology and Pathology of Platelet Vessel Wall
- 30 Interactions, "pp. 219-26 (1986), Crawford and Scrutton in: Bloom and Thomas, "Haemostasis and Thrombosis," pp. .47-77, (1987), Bevers, et. al., Eur. J. Biochem. 1982, 122, 429-36, Mann, Trends Biochem. Sci. 1987, 12, 229-33).
- 35 Thrombosis may be regarded as the pathological condition wherein improper activity of the hemostatic

mechanism results in intravascular thrombus formation. Etiological factors such as the presence of atherosclerotic plaque, phlebitis and septicemia may cause thrombosis, leading to impaired blood flow to the effected tissues and possible serious pathological consequences.

Currently, two of the most effective classes of drugs in clinical use as anticoagulants are the heparins and the vitamin K antagonists. The heparins are ill-defined mixtures of sulfated polysaccharides that bind to, and thus potentiate the action of antithrombin III. Antithrombin III is a naturally occurring inhibitor of the activated clotting factors IXa, Xa, XIa, thrombin and probably XIIa (see Jaques, Pharmacol. Rev. 1980, 31, pp. 99-166). The vitamin K antagonists, of which warfarin is the most well-known example, act indirectly by inhibiting the post-ribosomal carboxylations of the vitamin K dependent coagulation factors II, VII, IX and X (see Hirsch, Semin. Thromb. Hemostasis 1986, 12, 1-11). While effective therapies for the treatment of thrombosis, heparins and vitamin K antagonists have the unfortunate side effects of bleeding and marked interpatient variability, resulting in a small and unpredictable therapeutic safety margin. The use of direct acting thrombin inhibitors is expected to alleviate these problems.

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Thrombin is a serine protease having trypsin-like specificity for the cleavage of sequence-specific Arg-Xxx peptide bonds. As with other serine proteases, the cleavage event begins with an attack of the active site serine on the scissile bond of the substrate, resulting in the formation of a tetrahedral intermediate. This is followed by collapse of the tetrahedral intermediate to form an acyl enzyme and release of the amino terminus of the cleaved sequence. Hydrolysis of the acyl enzyme then releases the carboxy terminus.

A number of naturally occurring thrombin inhibitors have been reported. These include nazumamide A from Theonella sp. (see Fusetani, et. al., Tetrahedron Lett. 1991, 32, 7073-4), cyclotheonamide A from Theonella sp. (see Fusetani, et. al., J. Am. Chem. Soc. 1990, 112, 7053-4), amblyommin from Amblyomma hebraeum (see Bonin, et. al., EP 345614), hirudin from Hirudo medicinalis, recombinant versions of hirudin and hirudin fragments (see Rigbl and Jackson, EP 352903, Koerwer, WO 9109946, Meyer, et. al., WO 9108233, Dawson, et. al., WO 9109125, Maraganore, et. al., WO 9102750 and Maraganore, EP 333356).

Synthetic thrombin inhibitors have also been disclosed. Arylsulfonylarginine amides such as (2R, 4R)
4-methyl-1-[N^2 -{(3-methyl-1,2,3,4-tetrahydro-8-quinolinyl)sulfonyl}-L-arginyl]-2-piperidinecarboxylate have been shown to be effective inhibitors of thrombin (see Okamoto, et. al. Thromb Res. 1976, 8, 77-82, Ohshiro, et. al., Blood Vessel 1983, 14, 216-8), as have compounds containing constrained arginine mimics

- have compounds containing constrained arginine mimics such as (2-naphthylsulfonylglycyl)-4-amidino-phenylalanyl piperidide (see Stuerzebecher, et. al., Thromb. Res. 1983, 29, 635-42), 1-[2-[5-(dimethylamino)naphth-1-ylsulfonamido]-3-(2-
- iminohexahydropyrimidin-5-yl)propanoyl]-4methylpiperidine dihydrochloride (see Ishikawa, JP
 88227572 and Ishikawa and Inamura, JP 88227573), N(trans-4-amino-methylcyclohexylcarbonyl)-4-O-(2picolyl)-L-tyrosine 4-acetanilide dihydrochloride (see
- Okamoto, et. al., EP 217286) and 4[(aminoiminomethyl)amino]benzoic acid esters (see Fuji, et. al., DE 3005580, Matsuoka, et. al., Jpn. J. Pharmacol. 1989, 51, 455-63, and Takeshita, et. al., EP 435235).
- Inhibitor design has benefitted from the knowledge of the mechanism of action and of the peptide sequences

which are thought to bind in the catalytic site of thrombin, e.g., -Gly-Val-Arg-Gly- of fibrinogen (see Blombäck, et. al., J. Biol. Chem., 1972, 247, 1496-512), Ile-Pro-Arg-Ser- of prothrombin (see Magnussen, et. al., in: Reich, et. al., "Proteases and Biological Control, "pp. 123-149 (1975)) and -Val-Pro-Arg-Gly- of factor XIII (see Takagi and Doolittle, Biochemistry 1974, 13, 750-6 and Nakamura, et. al., Biochem. Biophys. Res. Commun. 1974, 58, 250-256). of mechanism-based inhibitors are exemplified by the tripeptide aldehyde D-Phe-Pro-N-Me-Arg-H (see Bajusz, et. al., J. Med. Chem. 1990, 33, 1729-35), the chloromethyl ketone Ac-(D)-Phe-Pro-ArgCH2Cl (see Kettner and Shaw, Thromb. Res. 1979, 14, 969-73) and the trifluoromethyl ketone D-Phe-Pro-ArgCF3 (see Kolb, et. 15 al., US 697987).

Kettner and Shenvi (EP 293881, published June 12, 1988), disclose peptide boronic acid inhibitors of trypsin-like proteases of formula (1)

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$$R^{1}-[(A^{3})_{q}(A^{2})_{p}(A^{1})_{o}]_{n}-NH-CHR^{2}-BY^{1}Y^{2}$$
 (1)

wherein Y^1 and Y^2 , independently, are hydroxyl or fluoro or, taken together, form a moiety derived from a dihydroxy compound having at least two hydroxy groups 25 separated by at least two connecting atoms in a chain or ring, said chain or ring comprising 1 to about 20 carbon atoms and, optionally, a heteroatom which can be N, S, or O; R² is a substituted alkyl selected from the group consisting of $-(CH_2)_z-X$, $-CH(CH_3)-(CH_2)_2-X$, $-CH_2-CH$ 30 $(CH_3) - CH_2 - X$, $-(CH_2)_2 - CH(CH_3) - X$ and $-(CH_2)_2 - CH(CH_3) - X$, where X is $-NH_2$, $-NH-C(NH)-NH_2$ or $-S-C(NH)-NH_2$, and z is 3 to 5; n, o, p and q are, independently, either 0 or 1; A^{1} , A^{2} and A^{3} are, independently, amino acids of L- or D-configuration selected from the group consisting of Ala, Arg, Asn, Asp, Cys, Gln, Glu, Gly, His, Ile, Leu, 35 Lys, Met, Phe, Pro, Ser, Thr, Trp, Tyr and Val; and R1

is a peptide comprised of 1 to about 20 amino acids, an acyl or a sulfonyl group comprised of 1 to about 20 carbon atoms, H, or an N-terminal protecting group. In this disclosure, Kettner and Shenvi demonstrated that the pinanediol esters of boropeptides are pharmacologically equivalent to the corresponding boronic acids.

Metternich (EP 0471651 A2) discloses borolysine thrombin inhibitors of formula (2)

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$W-Y-NR^4-CHR^5-BQ^1Q^2$ (2)

wherein W is an N-protecting group; Y is a sequence of n amino acids such that the n+1 amino acid peptide Y-Lys 15 or Y-Arg has an affinity for the active site of a trypsin-like protease; where n is an integer of from 1 to 10 and in which at least one amino acid is an unnatural amino acid having a hydrophobic side chain; O1 and Q^2 are the same or different and are selected from -OH, -COR1, -CONR1R2, -NR1R2 or -OR3 of Q^1 and Q^2 taken 20 together form a diol residue; R1, R2 and R3 which may be the same or different, are C1-10alkyl, C6-10aryl, C6-10aralkyl, or phenyl substituted by up to three groups selected from C₁₋₄alkyl, halogen and C₁₋₄alkoxy; R₄ is 25 hydrogen or C₁₋₁₀alkyl; R₅ is a group -A-X; wherein A is $-(CH_2)_z$ in which z is 2, 3, 4 or 5; $-CH(CH_3) - (CH_2)_2$; $-CH_2-CH(CH_3)-CH_2-$; $-(CH_2)_2-CH(CH_3)-$; $-(CH_2)_2-C(CH_3)_2-$; $CH(CH_3) - (CH_2)_3 - ; -CH_2 - CH(CH_3) - (CH_2)_2 - ; -CH_2 - CH_2 - CH(CH_3) - (CH_2)_3 - ; -CH_2 - CH_2 - CH$ CH_2- ; $-(CH_2)_3-CH(CH_3)-$; $-(CH_2)_3-C(CH_3)_2$: C_{6-10} aryl C_{6-10} 10aralkyl and X is -NH2, -NH-C(NH)-NH2, -S-C(NH)-NH2,-N3, 30 $-C_{1-4}$ alkoxy, C_{1-4} alkylthio or Si(CH₃)₃ or R₄ and R₅ taken together form a trimethylene group and the asymmetric carbon atom may have the D- or L-configuration or represent any mixture of these.

Surprising for their lack of a basic residue at P_1 are tripeptide thrombin inhibitors comprised of 1-

aminoboronic and 1-aminophosphonic acid analogs of 3-methoxy-propylglycine (see Claeson, et. al., US 07-245428) and pentylglycine (see Cheng, et. al., "Symposium on Thrombosis and Hemostasis," 1991, Amsterdam, Abstract 2150).

In addition to thrombin inhibition, boropeptides have been disclosed with utility as a treatment for tumors, viral infections and arthritis (US 4963655A and EP 354522A), emphysema (US 4499082A), hypertension (EP 315574A) and as factor VII/VIIa inhibitors (WO 8909612A). Kleemann, et. al. (AU A-24693/88) disclose renin-inhibiting 1-amino boronic acid derivatives of formula (3)

15 $A^{1}-A^{2}-HN-CHR^{2}-BXR^{3}(YR^{4})$ (3)

in which A^1 denotes a radical of formulae (4-8).

$$R^{1}NR^{6}-CHR^{5}-CO-$$

$$R^{1}CHR^{12}-CHR^{5}-CO-$$

$$R^{1}NR^{6}-CHR^{5}-CHR^{7}-CHR^{8}-CHR^{9}-CO-$$

$$R^{1}CHR^{12}-CHR^{5}-CHR^{7}-CHR^{8}-CHR^{9}-CO-$$

$$R^{10}-(CH_{2})_{n}-CH(CH_{2})_{m}R^{11}-CO-$$
(8)

Despite the foregoing, more efficacious and specific thrombin inhibitors are needed as potentially valuable therapeutic agents for the treatment of thrombosis.

None of the cited references describe or suggest the new thrombin-inhibiting boronic acid derivatives of the present invention.

Summary of Invention

The present invention pertains to novel compounds of formula (I):

compounds of formula (1)

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 $R^{1}-Z-CHR^{2}-BY^{1}Y^{2}$

(I)

wherein

 Y^1 and Y^2 are independently

- a) -OH,
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- b) -F,
- c) $-NR^3R^4$, or
- d) C1-C8-alkoxy;

 Y^1 and Y^2 when taken together can form

- a) a cyclic boron ester where said chain or ring contains from 2 to 20 carbon atoms and, optionally, a heteroatom which can be N, S, or O,
 - b) a divalent cyclic boro amide where said chain or ring contains from 2 to 20 carbon atoms,
 - c) a cyclic boro amide-ester where said chain or ring contains from 2 to 20 carbon atoms;

Z is

- a) $-(CH_2)_mCONR^8-$,
- b) -(CH₂)_mCSNR⁸-,
- c) -(CH₂)_mSO₂NR⁸-,
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- d) $-(CH_2)_mCO_2-$,
- e) $-(CH_2)_mC(S)O-$, or
- f) $-(CH_2)_mSO_2O_{-};$

 R^1 is

a) -(CH2)p-aryl, wherein aryl is phenyl, naphthyl or biphenyl substituted with one, two or three substituents selected from the group consisting of halo (F, Cl, Br, I), -CN, C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl, C2-C10-alkynyl, methylenedioxy, -R⁸, -OR⁸, -NO₂, -CF₃, -S(O)rR⁷,

- b) heteroaryl, wherein heteroaryl is an unsubstituted, monosubstituted or disubstituted:
- i) 5- or 6-membered aromatic ring, which contains from 1 to 3 heteroatoms selected from the group consisting of O, N, and S,
 - ii) quinolinyl,
 - iii) isoquinolinyl,
 - iv) benzopyranyl,
 - v) benzothiophenyl,
 - vi) benzofuranyl,
 - vii) 5,6,7,8-tetrahydroquinolinyl
 - viii) 5,6,7,8-tetrahydroisoquinolinyl

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and wherein the substituents are members selected from the group consisting of halo (F, Cl, Br, I), -CN, C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl, C2-C10-alkynyl, -R⁸, -OR⁸, -NO₂, -CF₃, -S(O)_TR⁷, -NR⁸R⁹, -COR⁸, -CO₂R⁸, -CONR⁸R⁹, NR⁸COR⁹, NRCO₂R⁹, NR¹²

c)

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d)

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e)

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f)

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 \mathbb{R}^2 is

'g)

- a) $-(CH_2)_n$ -NHC (NH) NH₂,
- b) $-(CH_2)_n$ -NHC (NH) NHCOCH3,

- c) $-(CH_2)_n-SC(NH)NH_2$,
- d) $-(CH_2)_n-SC(NH)NHCOCH_3$,
- e) $-(CH_2)_n-NH_2$, or
- f) $-(CH_2)_n-NH(2-pyridyl);$
- 5 R³ is H, phenyl or C1-C4-alkyl;

R⁴ is H or phenylsulfonyl;

 ${\tt R}^5$ and ${\tt R}^6$ are hydrogen or when taken together from a six membered aromatic ring optionally substituted with one, two or three substituents selected from the group

consisting of halo (F, C1, Br, I), -CN, C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl, C2-C10-alkynyl, $-OR^8$, $-NO_2$, $-CF_3$, $-S(O)_rR^7$, $-NR^8R^9$, $-COR^8$, $-CO_2R^8$, $-CONR^8R^9$, phenyl, benzyl, phenylethyl;

 R^7 is

- 15 a) phenyl,
 - b) C1-C4-alkyl,
 - c) C1-C4-alkoxy, or
 - d) -CF3;

R⁸ and R⁹ are independently

20 a) H,

b)

- c) C3-C7-cycloalkyl,
 - d) C1-C8-alkyl;

R¹⁰ and R¹¹ are independently

- a) halo (F, Cl, Br, I),
- b) -CN,
- 30 c) C1-C10-alkyl,
 - d) C3-C8-cycloalkyl,
 - e) C2-C10-alkenyl,
 - f) C2-C10-alkynyl,
 - $q) OR^8$

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h) -NO2,
          i) -CF3,
          j) -S(0) rR^{7}
          k) -NR^8R^9
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          1) -COR^9,
         m) - CO_2R^8
         n) -CONR^{8}R^{9};
     R^{12} is
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         a) H,
         b) C1-C4-alkyl,
         c) phenyl,
         d) benzyl
         e) -COR^7
         f) -SO_2R^7
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     m is 0 to 6;
     n is 3 or 4;
     p is 0 to 2;
     r is 0 to 2;
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     t is 1 to 5
     E is -CO-, -SO<sub>2</sub>-, -CH<sub>2</sub>- or a single bond,
     F is -CO-; and pharmaceutically acceptable salts
     thereof.
          Preferred compounds of formula (I) are those
     compounds wherein R1 is phenyl and biphenyl containing
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     1-3 substituents selected from the series halo (F, Cl,
     Br, I), C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl,
     C2-C10-alkynyl, -R^8, -OR^8, -NO_2, -CF_3, -S(O)_rR^7, -NR^8R^9,
     -COR8, -CO2R8, -CONR8R9; NR8COR9;
    \mathbb{R}^2 is
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More preferred are those preferred compounds wherein $z = -(CH_2)_m CONR^8-$.

a) $-(CH_2)_3-NHC(NH)NH_2$, or

b) $-(CH_2)_3-SC(NH)NH_2$.

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Most preferred are those more preferred compounds listed below: N^{1} -(4-phenylbenzoyl)-(R)-boroarginine, hydrochloride N^{1} -(3-phenoxybenzoyl)-(R)-boroarginine, hydrochloride N^{1} -(1-fluorenonyl)-(R)-boroarginine, hydrochloride N^{1} -(4-[1-butyl]benzoyl)-(R)-boroarginine, hydrochloride N^{1} -(2-benzoylbenzoyl)-(R)-boroarginine, hydrochloride N^{1} -(5-phenyl-2-furoyl)-(R)-boroarginine, hydrochloride N^{1} -(3-[N-benzyloxycarbonyl-N-methylamino]-4-[1-butyl]benzoyl) - (R) -boroarginine, hydrochloride N^{1} -(2-phenyl-4-isoquinolyl)-(R)-boroarginine, hydrochloride · N^{1} -(4-cyclohexylbenzoyl)-(R)-boroarginine, hydrochloride 15 N^{1} -(2-methyl-4-phenylbenzoyl)-(R)-boroarginine, hydrochloride -Illustrative of the compounds of this invention are the following: 20 N^{1} -(4-phenylbenzoyl)-(R)-boroarginine (+)-pinanediol, bisulfite N^{1} -(3-phenylbenzoyl)-(R)-boroarginine (+)-pinanediol, bisulfite 25 N^{1} -(3-phenoxybenzoyl)-(R)-boroarginine (+)-pinanediol, bisulfite N^{1} -(4-[4-pyridyl]benzoyl)-(R)-boroarginine (+)pinanediol, bisulfite N^{1} -(2-benzoylbenzoyl)-(R)-boroarginine (+)-pinanediol. bisulfite N^{1} -(3-benzoylbenzoyl)-(R)-boroarginine (+)-pinanediol, bisulfite

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bisulfite

 N^{1} -(4-benzoylbenzoyl)-(R)-boroarginine (+)-pinanediol,

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N^{1}-(3-[N-benzyloxycarbonyl]aminobenzoyl)-(R)-
     boroarginine (+)-pinanediol, bisulfite
     N^{1}-(3-[N-benzyloxycarbonyl-N-methyl]aminobenzoyl)-(R)-
     boroarginine (+)-pinanediol, bisulfite
     N^{1}-(4-ethylbenzoyl)-(R)-boroarginine (+)-pinanediol,
     bisulfite
     N^{1}-(4-n-propylbenzoyl)-(R)-boroarginine (+)-pinanediol,
     bisulfite
     N^{1}-(4-isopropylbenzoyl)-(R)-boroarginine (+)-pinanediol,
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     bisulfite
     N^{1}-(4-n-butylbenzoyl)-(R)-boroarginine (+)-pinanediol,
     bisulfite
     N^{1}-(4-tert-butylbenzoyl)-(R)-boroarginine (+)-
     pinanediol, bisulfite
     N^{1}-(4-n-hexylbenzoyl)-(R)-boroarginine (+)-pinanediol,
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     bisulfite
     N^{1}-(4-cyclohexylbenzoyl)-(R)-boroarginine (+)-
     pinanediol, bisulfite
     N^{1}-(2-[N-(2-phenylethyl)carbonyl]aminobenzoyl)-(R)-
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     boroarginine (+)-pinanediol, bisulfite
     N^{1}-(4-n-butyloxybenzoyl)-(R)-boroarginine (+)-
    pinanediol, bisulfite
    N^{1}-(4-[N-cyclopropylcarbonyl]aminobenzoyl)-(R)-
    boroarginine (+)-pinanediol, bisulfite
25
    N^{1} (4-[N-cyclohexylcarbonyl] aminobenzoyl) - (R) -
    boroarginine (+)-pinanediol, bisulfite
     N^{1}-(4-[N-(4-methoxy)benzoyl]aminobenzoyl)-(R)-
    boroarginine (+)-pinanediol, bisulfite
    N^{1}-(4-[4-methoxy]phenylbenzoyl)-(R)-boroarginine (+)-
    pinanediol, bisulfite
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    N^{1}-(2-[2-phenyl]benzyloxycarbonylbenzoyl)-(R)-
    boroarginine (+)-pinanediol, bisulfite
    N^{1}-(2-[1-naphthyl]benzoyl)-(R)-boroarginine (+)-
    pinanediol, bisulfite
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    N^{1}-(4-[4-carboxy]phenylbenzoyl)-(R)-boroarginine (+)-
    pinanediol,
                  bisulfite
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N^{1}-(4-phenylbenzoyl)-(R)-borothioarginine (+)-
     pinanediol, hydrobromide
     N^{1}-(3-phenylbenzoyl)-(R)-borothioarginine (+)-
     pinanediol, hydrobromide
     N^{1}-(3-phenoxybenzoyl)-(R)-borothioarginine (+)-
     pinanediol,
                  hydrobromide
     N^{1}-(2-benzoylbenzoyl)-(R)-borothioarginine (+)-
     pinanediol, hydrobromide
     N^{1}-(3-benzoylbenzoyl)-(R)-borothioarginine (+)-
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     pinanediol, hydrobromide
     N^{1}-(4-benzoylbenzoyl)-(R)-borothioarginine (+)-
     pinanediol, hydrobromide
     N^{1}-(3-[N-benzyloxycarbonyl]aminobenzoyl)-(R)-
     borothioarginine (+)-pinanediol, hydrobromide
     N^{1}-(3-[N-benzyloxycarbonyl-N-methyl]aminobenzoyl)-(R)-
     borothioarginine (+)-pinanediol, hydrobromide
     N^{1}-(4-ethylbenzoyl)-(R)-borothioarginine (+)-pinanediol,
     hydrobromide
     N^{1}-(4-n-propylbenzoyl)-(R)-borothioarginine (+)-
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    pinanediol, hydrobromide
     N^{1}-(4-isopropylbenzoyl)-(R)-borothioarginine (+)-
    pinanediol, hydrobromide
     N^{1}-(4-n-butylbenzoyl)-(R)-borothioarginine (+)-
    pinanediol, hydrobromide
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    N^{1}-(4-tert-butylbenzoyl)-(R)-borothioarginine (+)-
    pinanediol,
                  hydrobromide
    N^{1}-(4-n-hexylbenzoyl)-(R)-borothioarginine (+)-
    pinanediol, hydrobromide
    N^{1}-(4-cyclohexylbenzoyl)-(R)-borothioarginine (+)-
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    pinanediol, hydrobromide
    N^{1}-(2-[N-(2-phenylethyl)carbonyl]aminobenzoyl)-(R)-
    borothioarginine (+)-pinanediol, hydrobromide
    N^{1}-(4-n-butyloxybenzoyl)-(R)-borothioarginine (+)-
    pinanediol, hydrobromide
35
    N^{1}- (4-[N-cyclopropylcarbonyl] aminobenzoyl) - (R) -
   borothioarginine (+)-pinanediol, hydrobromide
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 N^{1} -(4-[N-cyclohexylcarbonyl]aminobenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(4-[N-(4-methoxy)benzoyl]aminobenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(4-[4-methoxy]phenylbenzoyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide N^{1} (2-[2-phenylbenzyloxycarbonyl]benzoyl) - (R) borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-[1-naphthyl]benzoyl)-(R)-borothioarginine (+)pinanediol, hydrobromide N^{1} -(4-[4-carboxy]phenylbenzoyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide N^{1} -([2-anthraquinonyl]carbonyl)-(R)-boroarginine (+)pinanediol, bisulfite N^{1} -([2-dioxothioxanthinonyl]carbonyl)-(R)-boroarginine 15 (+)-pinanediol, bisulfite N^{1} -([2-anthraquinonyl]carbonyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide N^{1} -([2-dioxothioxanthinonyl]carbonyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -([2-fluoren-9-onyl]carbonyl)-(R)-borothiohomoarginine (+)-pinanediol, hydrobromide N^{1} -([2-fluoren-9-onyl]carbonyl)-(R)-boroarginine (+)pinanediol, bisulfite N^{2} -([2-fluoren-9-onyl]carbonyl)-(R)-borothioarginine 25 (+)-pinanediol, hydrobromide N^{2} -([3-fluoren-9-onyl]carbonyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide N^{1} -([3-fluoren-9-onyl]carbonyl)-(R)-boroarginine (+)-- 30 pinanediol, bisulfite N^{1} -([4-fluoren-9-onyl]carbonyl)-(R)-borothioarginine

 N^{1} -([4-fluoren-9-onyl]carbonyl)-(R)-boroarginine (+)-

 N^{1} -(1-naphthoy1)-(R)-borothioarginine (+)-pinanediol,

(+)-pinanediol, hydrobromide

pinanediol, bisulfite

hydrobromide

 N^{1} -(1-naphthoyl)-(R)-boroarginine (+)-pinanediol, bisulfite N^{1} (2-methyl-4-phenyl-5-methoxybenzoyl) - (R) borothioarginine (+)-pinanediol, hydrobromide 5 N^{1} -(2-methyl-4-phenyl-5-carboxamidobenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-phenyl-5-fluorobenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-phenyl-5-trifluoromethylbenzoyl)-(R)borothioarginine (+)-pinanediol, 10 hydrobromide N^{1} -(2-methyl-4-phenyl-5-chlorobenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-phenyl-5-hydroxybenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-[4-carboxy]phenyl-5-methoxybenzoyl)-(R)-15 borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-[4-carboxy]phenyl-5-carboxamidobenzoyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-[4-carboxy]phenyl-5-fluorobenzoyl)-(R)-20 borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-[4-carboxy]phenyl-5trifluoromethylbenzoyl) - (R) - borothioarginine (+) pinanediol, hydrobromide N^{1} -(2-methyl-4-[4-carboxy]phenyl-5-chlorobenzoyl)-(R)-25 borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-[4-carboxy]phenyl-5-hydroxybenzoyl)-(R)borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-methyl-4-phenyl-5-methoxybenzoyl)-(R)-boroarginine (+)-pinanediol, bisulfite 30 N^{1} -(2-methyl-4-phenyl-5-carboxamidobenzoyl)-(R)boroarginine (+)-pinanediol, bisulfite N^{1} -(2-methyl-4-phenyl-5-fluorobenzoyl)-(R)-boroarginine (+)-pinanediol, bisulfite N^{1} -(2-methyl-4-phenyl-5-trifluoromethylbenzoyl)-(R)-35 boroarginine (+)-pinanediol, bisulfite

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N^{1}-(2-methyl-4-phenyl-5-chlorobenzoyl)-(R)-boroarginine
     (+)-pinanediol, bisulfite
     N^{1}-(2-methyl-4-phenyl-5-hydroxybenzoyl)-(R)-boroarginine
     (+)-pinanediol, bisulfite
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-methoxybenzoyl)-(R)-
     boroarginine (+)-pinanediol, bisulfite
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-carboxamidobenzoyl)-
     (R)-boroarginine (+)-pinanediol, bisulfite
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-fluorobenzoyl)-(R)-
     boroarginine (+)-pinanediol, bisulfite
10
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-
     trifluoromethylbenzoyl) - (R) -boroarginine (+) -pinanediol,
     bisulfite
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-chlorobenzoyl)-(R)-
     boroarginine (+)-pinanediol, bisulfite
15
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-hydroxybenzoyl)-(R)-
     boroarginine (+)-pinanediol, bisulfite
     N^{1}-(2-[5-phenyl]furylcarbonyl)-(R)-boroarginine (+)-
     pinanediol, bisulfite
20
     N^{2}-(2-[5-phenyl]thiophenylcarbonyl)-(R)-boroarginine
     (+)-pinanediol, bisulfite
     N^{1}-(2-[5-phenyl]furylcarbonyl)-(R)-borothioarginine (+)-
     pinanediol, hydrobromide
     N^{1}-(2-[5-phenyl]thiophen-ylcarbonyl)-(R)-
25
    borothioarginine (+)-pinanediol, hydrobromide
     N^{1}-(3-[6-phenyl]pyridylcarbonyl)-(R)-boroarginine (+)-
    pinanediol, bisulfite
     N^{1}-(3-[5-benzyloxy]pyridylcarbonyl)-(R)-boroarginine
     (+)-pinanediol, bisulfite
    N^{1}-(3-[6-phenyl]pyridylcarbonyl)-(R)-borothioarginine
30
     (+)-pinanediol, hydrobromide
    N^{1}-(3-[5-benzyloxy]pyridylcarbonyl)-(R)-borothioarginine
    ·(+)-pinanediol, hydrobromide
    N^{1}-(2-benzopyronylcarbonyl)-(R)-boroarginine (+)-
    pinanediol, bisulfite
35
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(%)

 N^{1} -(2-benzopyronylcarbonyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide N^{1} -(3-isoquinolinylcarbonyl)-(R)-boroarginine (+)-pinanediol, bisulfite

- 5 N¹-(2-phenyl-4-isoquinolinylcarbonyl)-(R)-boroarginine (+)-pinanediol, bisulfite N¹-(3-isoquinolinylcarbonyl)-(R)-borothioarginine (+)pinanediol, hydrobromide N¹-(2-phenyl-4-isoquinolinylcarbonyl)-(R)-
- borothioarginine (+)-pinanediol, hydrobromide N^{1} -(2-isoquinolinylcarbonyl)-(R)-boroarginine (+)-pinanediol, bisulfite N^{1} -(2-isoquinolinylcarbonyl)-(R)-borothioarginine (+)-pinanediol, hydrobromide

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- N¹-(4-phenylbenzoyl)-(R)-boroarginine, hydrochloride N¹-(3-phenylbenzoyl)-(R)-boroarginine, hydrochloride N¹-(3-phenoxybenzoyl)-(R)-boroarginine, hydrochloride N¹-(4-[4-pyridyl]benzoyl)-(R)-boroarginine, hydrochloride
- 20 N^{1} -(2-benzoylbenzoyl)-(R)-boroarginine, hydrochloride N^{1} -(3-benzoylbenzoyl)-(R)-boroarginine, hydrochloride N^{1} -(4-benzoylbenzoyl)-(R)-boroarginine, hydrochloride N^{1} -(3-[N-benzyloxycarbonyl]aminobenzoyl)-(R)-boroarginine, hydrochloride
- N¹-(3-[N-benzyloxycarbonyl-N-methyl]aminobenzoyl)-(R)-boroarginine, hydrochloride
 N¹-(4-ethylbenzoyl)-(R)-boroarginine, hydrochloride
 N¹-(4-n-propylbenzoyl)-(R)-boroarginine, hydrochloride
 N¹-(4-isopropylbenzoyl)-(R)-boroarginine, hydrochloride
- 30 N¹-(4-tert-butylbenzoyl)-(R)-boroarginine,
 hydrochloride
 N¹-(4-n-hexylbenzoyl)-(R)-boroarginine, hydrochloride
 N¹-(4-cyclohexylbenzoyl)-(R)-boroarginine,
 hydrochloride
- 35 N^{1} -(2-[N-(2-phenylethyl)carbonyl]aminobenzoyl)-(R)-boroarginine, hydrochloride

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N^{1}-(4-n-butyloxybenzoyl)-(R)-boroarginine,
     hydrochloride
     N^{1}-(4-[N-cyclopropylcarbonyl]aminobenzoyl)-(R)-
     boroarginine, hydrochloride
 5 N^{1}-(4-[N-cyclohexylcarbonyl]aminobenzoyl)-(R)-
    boroarginine, hydrochloride
     N^{1}-(4-[N-(4-methoxy)benzoyl]aminobenzoyl)-(R)-
    boroarginine, hydrochloride
    N^{1}-(4-[4-methoxy]phenylbenzoyl)-(R)-boroarginine,
10
    hydrochloride
    N^{1}-(2-[2-phenyl]benzyloxycarbonylbenzoyl)-(R)-
    boroarginine, hydrochloride
    N^{1}-(2-[1-naphthyl]benzoyl)-(R)-boroarginine,
    hydrochloride
    N^{1}-(4-[4-carboxy]phenylbenzoyl)-(R)-boroarginine,
15
    hydrochloride
    N^{1}-([2-anthraquinonyl]carbonyl)-(R)-boroarginine,
    hydrochloride
    N^{1}-([2-dioxothioxanthinonyl]carbonyl)-(R)-boroarginine,
20
    hydrochloride
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hydrochloride

25 N¹-(1-naphthoy1)-(R)-boroarginine, hydrochloride
N¹-([4-fluoren-9-onyl]carbonyl)-(R)-boroarginine,
hydrochloride

 N^{1} -([2-fluoren-9-onyl]carbonyl)-(R)-boroarginine,

 N^{1} -([3-fluoren-9-onyl]carbonyl)-(R)-boroarginine,

- N^{1} -(2-methyl-4-phenyl-5-methoxybenzoyl)-(R)-boroarginine, hydrochloride
- 30 N¹-(2-methyl-4-phenyl-5-carboxamidobenzoyl)-(R)boroarginine, hydrochloride
 N¹-(2-methyl-4-phenyl-5-fluorobenzoyl)-(R)-boroarginine,
 hydrochloride
 N¹-(2-methyl-4-phenyl-5-trifluoromethylbenzoyl)-(R)-
- 35 boroarginine, hydrochloride

hydrochloride

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N^{2}-(2-methyl-4-phenyl-5-chlorobenzoyl)-(R)-boroarginine,
     hydrochloride
     N^{1}-(2-methyl-4-phenyl-5-hydroxybenzoyl)-(R)-
     boroarginine, hydrochloride
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-methoxybenzoyl)-(R)-
     boroarginine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-carboxamidobenzoyl)-
      (R)-boroarginine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-fluorobenzoyl)-(R)-
    boroarginine, hydrochloride
10
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-
     trifluoromethylbenzoyl)-(R)-boroarginine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-chlorobenzoyl)-(R)-
     boroarginine, hydrochloride
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-hydroxybenzoyl)-(R)-
15
     boroarginine, hydrochloride
     N^{1}-(2-[5-phenyl]furylcarbonyl)-(R)-boroarginine,
     hydrochloride
     N^{1}-(2-[5-phenyl]thiophenylcarbonyl)-(R)-boroarginine,
20
     hydrochloride
     N^{1}-(2-benzopyronylcarbonyl)-(R)-boroarginine,
     hydrochloride
     N^{1}-(2-isoquinolinylcarbonyl)-(R)-boroarginine,
    hydrochloride
25
    N^{1}-(3-isoquinolinylcarbonyl)-(R)-boroarginine,
    hydrochloride
     N^{1}-(2-phenyl-4-isoquinolinylcarbonyl)-(R)-boroarginine,
    hydrochloride
    N^{1}-(4-phenylbenzoyl)-(R)-borothioarginine,
30
    hydrochloride
    N^{1}-(3-phenylbenzoyl)-(R)-borothioarginine,
    hydrochloride
    N^{1}-(3-phenoxybenzoyl)-(R)-borothioarginine,
    hydrochloride
    N^{1}-(2-benzoylbenzoyl)-(R)-borothioarginine,
35
    hvdrochloride
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N^{1}-(3-benzoylbenzoyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(4-benzoylbenzoyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(3-[N-benzyloxycarbonyl]aminobenzoyl)-(R)-
     borothioarginine, hydrochloride
     N^{1}-(3-[N-benzyloxycarbonyl-N-methyl]aminobenzoyl)-(R)-
     borothioarginine, hydrochloride
     N^{1}-(4-ethylbenzoyl)-(R)-borothioarginine, hydrochloride
     N^{1}- (4-n-propylbenzoyl) - (R)-borothioarginine,
     hydrochloride
     N^{2}-(4-isopropylbenzoyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(4-n-butylbenzoyl)-(R)-borothioarginine,
15
     hydrochloride
     N^{1}-(4-tert-butylbenzoyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(4-n-hexylbenzoyl)-(R)-borothioarginine,
     hydrochloride
20
     N^{1}- (4-cyclohexylbenzoyl) - (R)-borothioarginine,
     hydrochloride
     N^{1}-(2-[N-(2-phenylethyl)carbonyl]aminobenzoyl)-(R)-
     borothioarginine, hydrochloride
     N^{1}-(4-n-butyloxybenzoyl)-(R)-borothioarginine,
   hydrochloride
     N^{1}-(4-[N-cyclopropylcarbonyl]aminobenzoyl)-(R)-
     borothioarginine, hydrochloride
     N^{1}- (4-[N-cyclohexylcarbonyl]aminobenzoyl) - (R) -
    borothioarginine,
                         hydrochloride
30 N^{1}-(4-[N-(4-methoxy)benzoyl]aminobenzoyl)-(R)-
    borothioarginine, hydrochloride
    N^{1}-(4-[4-methoxy]phenylbenzoyl)-(R)-borothioarginine,
    hydrochloride
    N^{1}-(2-[2-phenylbenzyloxycarbonyl]benzoyl)-(R)-
35
    borothioarginine, hydrochloride
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N^{1}-(2-[1-naphthyl]benzoyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(4-[4-carboxy]phenylbenzoyl)-(R)-borothioarginine,
     hydrochloride
 5
     N^{1}-([2-anthraquinonyl]carbonyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-([2-dioxothioxanthinonyl]carbonyl)-(R)-
     borothioarginine, hydrochloride
     N^{1}-([2-fluoren-9-onyl]carbonyl)-(R)-
     borothiohomoarginine, hydrochloride
     N^{1}-([2-fluoren-9-onyl]carbonyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-([3-fluoren-9-onyl]carbonyl)-(R)-borothioarginine,
     hvdrochloride
     N^{1}-([4-fluoren-9-onyl]carbonyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(1-naphthoyl)-(R)-borothioarginine, hydrochloride
     N^{1}-(2-methyl-4-phenyl-5-methoxybenzoyl)-(R)-
    borothioarginine, hydrochloride
20
    N^{1}-(2-methyl-4-phenyl-5-carboxamidobenzoyl)-(R)-
    borothioarginine, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-fluorobenzoyl)-(R)-
    borothioarginine, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-trifluoromethylbenzoyl)-(R)-
    borothioarginine, hydrochloride
25
    N^{1}-(2-methyl-4-phenyl-5-chlorobenzoyl)-(R)-
    borothioarginine, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-hydroxybenzoyl)-(R)-
    borothioarginine, hydrochloride
    N^{2}-(2-methyl-4-[4-carboxy]phenyl-5-methoxybenzoyl)-(R)-
30
    borothioarginine, hydrochloride
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-carboxamidobenzoyl)-
     (R)-borothioarginine, hydrochloride
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-fluorobenzoyl)-(R)-
    borothioarginine, hydrochloride
35
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N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-
     trifluoromethylbenzoyl) - (R) -borothioarginine,
     hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-chlorobenzovl)-(R)-
     borothioarginine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-hydroxybenzoyl)-(R)-
     borothioarginine,
                         hydrochloride
     N^{1}-(2-[5-phenyl]furylcarbonyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(2-[5-phenyl]thiophenylcarbonyl)-(R)-
     borothioarginine, hydrochloride
     N^{1}-(3-[6-phenyl]pyridylcarbonyl)-(R)-boroarginine,
     hydrochloride
     N^{1}-(3-[5-benzyloxy]pyridylcarbonyl)-(R)-boroarginine,
15
    hydrochloride
     N^{1}-(3-[6-phenyl]pyridylcarbonyl)-(R)-borothioarginine,
     hydrochloride
     N^{1}-(3-[5-benzyloxy]pyridylcarbonyl)-(R)-
    borothioarginine, hydrochloride
20
    N^{1} (2-benzopyronylcarbonyl) - (R)-borothioarginine,
    hydrochloride
     N^{1}-(3-isoquinolinylcarbonyl)-(R)-borothioarginine,
    hydrochloride
    N^{1}-(2-phenyl-4-isoquinolinylcarbonyl)-(R)-
    borothioarginine, hydrochloride
25
    N^{1}-(2-isoquinolinylcarbonyl)-(R)-borothioarginine,
    hydrochloride
    N^{1}-(4-phenylbenzoyl)-(R)-borolysine (+)-pinanediol,
    hydrochloride
30
    N^{1}-(3-phenylbenzoyl)-(R)-borolysine (+)-pinanediol,
    hydrochloride
    N^{2}-(3-phenoxybenzoyl)-(R)-borolysine (+)-pinanediol,
    hydrochloride
    N^{2}-(4-[4-pyridyl]benzoyl)-(R)-borolysine (+)-pinanediol,
35
    hydrochloride
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N^{1}-(2-benzoylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
     N^{1}-(3-benzoylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
     N^{1}-(4-benzoylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
     N^{1}-(3-[N-benzyloxycarbonyl]aminobenzoyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
     N^{1}-(3-[N-benzyloxycarbonyl-N-methyl]aminobenzoyl)-(R)-
10
     borolysine (+)-pinanediol, hydrochloride
     N^{1}-(4-ethylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
     N^{1}-(4-n-propylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
15 N^{1}-(4-isopropylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
     N^{1}-(4-tert-butylbenzoyl)-(R)-borolysine (+)-pinanediol,
     hydrochloride
     N^{1}-(4-n-hexylbenzoyl)-(R)-borolysine (+)-pinanediol,
20
    hydrochloride
     N^{1}-(4-cyclohexylbenzoyl)-(R)-borolysine (+)-pinanediol,
    hydrochloride
    N^{1}-(2-[N-(2-phenylethyl) carbonyl] aminobenzoyl) - (R) -
    borolysine (+)-pinanediol, hydrochloride
25
    N^{1}-(4-n-butyloxybenzoyl)-(R)-borolysine (+)-pinanediol,
    hydrochloride
    N^{1}-(4-[N-cyclopropylcarbonyl]aminobenzoyl)-(R)-
    borolysine (+)-pinanediol, hydrochloride
    N^{1}-(4-[N-cyclohexylcarbonyl]aminobenzoyl)-(R)-borolysine
30
     (+)-pinanediol, hydrochloride
    N^{1}-(4-[N-(4-methoxy)benzoyl]aminobenzoyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
    N^{2}-(4-[4-methoxy]phenylbenzoyl)-(R)-borolysine (+)-
    pinanediol, hydrochloride
35
    N^{1}-(2-[2-phenyl]benzyloxycarbonylbenzoyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
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N^{1}-(2-[1-naphthyl]benzoyl)-(R)-borolysine (+)-
     pinanediol, hydrochloride
     N^{1}-(4-[4-carboxy]phenylbenzoyl)-(R)-borolysine (+)-
     pinanediol, hydrochloride
     N^{1}-([2-anthraquinonyl]carbonyl)-(R)-borolysine (+)-
    pinanediol, hydrochloride
     N^{1}-([2-dioxothioxanthinonyl]carbonyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
     N^{1}-([2-fluoren-9-onyl]carbonyl)-(R)-borolysine (+)-
10
    pinanediol, hydrochloride
     N^{1}-([3-fluoren-9-onyl]carbonyl)-(R)-borolysine (+)-
    pinanediol, hydrochloride
    N^{1}-(1-naphthoyl)-(R)-borolysine (+)-pinanediol,
    hydrochloride
15
    N^{1}-([4-fluoren-9-onyl]carbonyl)-(R)-borolysine (+)-
    pinanediol, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-methoxybenzoyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-carboxamidobenzoyl)-(R)-
20
    borolysine (+)-pinanediol, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-fluorobenzoyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-trifluoromethylbenzoyl)-(R)-
    borolysine (+)-pinanediol, hydrochloride
   N^{1}-(2-methyl-4-phenyl-5-chlorobenzoyl)-(R)-borolysine
    (+)-pinanediol, hydrochloride
    N^{2}-(2-methyl-4-phenyl-5-hydroxybenzoyl)-(R)-borolysine
    (+)-pinanediol, hydrochloride
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-methoxybenzoyl)-(R)-
30
    borolysine (+)-pinanediol, hydrochloride
    N<sup>1</sup>-(2-methyl-4-[4-carboxy]phenyl-5-carboxamidobenzoyl)-
    (R)-borolysine (+)-pinanediol, hydrochloride
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-fluorobenzoyl)-(R)-
    borolysine (+)-pinanediol, hydrochloride
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N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-
      trifluoromethylbenzoyl) - (R) -borolysine (+) -pinanediol,
      hydrochloride
      N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-chlorobenzoyl)-(R)-
     borolysine (+)-pinanediol, hydrochloride
      N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-hydroxybenzoyl)-(R)-
     borolysine (+)-pinanediol, hydrochloride
     N^{1}-(2-[5-phenyl]furylcarbonyl)-(R)-borolysine (+)-
     pinanediol, hydrochloride
     N^{1}-(2-[5-phenyl]thiophen-ylcarbonyl)-(R)-borolysine (+)-
 10
     pinanediol, hydrochloride
     N^{1}-(2-benzopyronylcarbonyl)-(R)-borolysine (+)-
     pinanediol, hydrochloride
     N^{1}-(2-isoquinolinylcarbonyl)-(R)-borolysine (+)-
     pinanediol, hydrochloride
15
     N^{1}-(3-isoquinolinylcarbonyl)-(R)-borolysine (+)-
     pinanediol, hydrochloride
     N^{1}-(2-phenyl-4-isoquinolinylcarbonyl)-(R)-borolysine
     (+)-pinanediol, hydrochloride
20
     N^{1}-(4-phenylbenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(3-phenylbenzoyl)-(R)-borolysine, hydrochloride
    N^{1}-(3-phenoxybenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(4-[4-pyridyl]benzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(2-benzoylbenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(3-benzoylbenzoyl)-(R)-borolysine, hydrochloride
25
     N^{1}-(4-benzoylbenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(3-[N-benzyloxycarbonyl]aminobenzoyl)-(R)-borolysine,
     hydrochloride
     N^{1}-(3-[N-benzyloxycarbonyl-N-methyl]aminobenzoyl)-(R)-
30
     borolysine, hydrochloride
     N^{1}-(4-ethylbenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(4-n-propylbenzoyl)-(R)-borolysine,
                                              hydrochloride
     N^{1}-(4-isopropylbenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(4-tert-butylbenzoyl)-(R)-borolysine, hydrochloride
     N^{1}-(4-n-hexylbenzoyl)-(R)-borolysine, hydrochloride
35
     N^{1}-(4-cyclohexylbenzoyl)-(R)-borolysine, hydrochloride
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N^{1} (2-[N-(2-phenylethyl) carbonyl] aminobenzoyl) - (R) -
     borolysine, hydrochloride
     N^{2}-(4-n-butyloxybenzoyl)-(R)-borolysine, hydrochloride
     N^{1} - (4-[N-cyclopropylcarbonyl] aminobenzoyl) - (R) -
 5 borolysine, hydrochloride
     N^{1} - (4-[N-cyclohexylcarbonyl]aminobenzoyl) - (R) -
     borolysine, hydrochloride
     N^{1}-(4-[N-(4-methoxy)benzoyl]aminobenzoyl)-(R)-
     borolysine, hydrochloride
10
     N^{1}- (4-[4-methoxy]phenylbenzoyl) - (R) -borolysine,
     hydrochloride
     N^{1}-(2-[2-phenyl]benzyloxycarbonylbenzoyl)-(R)-
    borolysine, hydrochloride
    N^{1}-(2-[1-naphthyl]benzoyl)-(R)-borolysine,
15
    hydrochloride
    N^{1}-(4-[4-carboxy]phenylbenzoyl)-(R)-borolysine,
     hydrochloride
     N^{1}-([2-anthraquinonyl]carbonyl)-(R)-borolysine,
    hydrochloride
20
    N^{1}-([2-dioxothioxanthinonyl]carbonyl)-(R)-borolysine,
     hydrochloride
     N^{1}-([2-fluoren-9-onyl]carbonyl)-(R)-borolysine,
    hydrochloride
     N^{2}-([3-fluoren-9-onyl]carbonyl)-(R)-borolysine,
25 hydrochloride
    N^{2}-(1-naphthoy1)-(R)-borolysine, hydrochloride
    N^{1}-([4-fluoren-9-onyl]carbonyl)-(R)-borolysine,
    hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-methoxybenzoyl)-(R)-borolysine,
30
    hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-carboxamidobenzoyl)-(R)-
    borolysine, hydrochloride
    N^{1}-(2-methyl-4-phenyl-5-fluorobenzoyl)-(R)-borolysine,
    hydrochloride
35
   N^{2} (2-methyl-4-phenyl-5-trifluoromethylbenzoyl) - (R) -
    borolysine, hydrochloride
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N^{1}-(2-methyl-4-phenyl-5-chlorobenzoyl)-(R)-borolysine,
     hydrochloride
     N^{1}-(2-methyl-4-phenyl-5-hydroxybenzoyl)-(R)-borolysine,
     hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-methoxybenzoyl)-(R)-
     borolysine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-carboxamidobenzoyl)-
     (R)-borolysine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-fluorobenzoyl)-(R)-
    borolysine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-
     trifluoromethylbenzoyl) - (R) -borolysine, hydrochloride
     N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-chlorobenzoyl)-(R)-
    borolysine, hydrochloride
15
    N^{1}-(2-methyl-4-[4-carboxy]phenyl-5-hydroxybenzoyl)-(R)-
    borolysine, hydrochloride
    N^{1}-(2-[5-phenyl]furylcarbonyl)-(R)-borolysine,
    hydrochloride
    N^{1}-(2-[5-phenyl]thiophen-ylcarbonyl)-(R)-borolysine,
20
    hydrochloride
    N^{1}-(2-benzopyronylcarbonyl)-(R)-borolysine,
    hydrochloride
    N^{1}-(2-isoquinolinylcarbonyl)-(R)-borolysine,
    hydrochloride
    N^{1}-(3-isoquinolinylcarbonyl)-(R)-borolysine,
25
    hydrochloride
    N^{2}-(2-phenyl-4-isoquinolinylcarbonyl)-(R)-borolysine,
    hydrochloride
    N^{2}-(2-methyl-4-phenylbenzoyl)-R-borolysine,
30
    hydrochloride
    N^{1}-(2-methyl-4-phenylbenzoyl)-R-borolysine, (+)-
    pinanediol, hydrochloride
    N^{1}-(2-methyl-4-phenylbenzoyl)-R-borothioarginine (+)-
    hydrobromide
    N^{1}-(2-methyl-4-phenylbenzoyl)-R-borothioarginine (+)-
35
    pinanediol, hydrochloride
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 N^{1} -(2-methyl-4-phenylbenzoyl)-R-boroarginine (+)-hydrochloride N^{1} -(2-methyl-4-phenylbenzoyl)-R-boroarginine (+)-pinanediol, bisulfite

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Detailed Description of the Invention

Throughout the specification, the following conventional three-letter abbreviations for amino acid residues or amino acids apply:

Ala = alanine

Arg = arginine

Asn = asparagine

Asp = aspartic acid

Cys = cysteine

Gln = glutamine

Glu = glutamic acid

Gly = glycine

His = histidine

20 Ile = isoleucine

Leu = leucine

Lys = lysine

Met = methionine

Phe = phenylalanine

25 Pro = proline

Ser = serine

Thr = threonine

Trp = tryptophan

Tyr = tyrosine

30 Val = valine

The prefix "boro" indicates amino acid residues where the carboxy group is replaced by a boronic acid (Formula I, Y^1 and $Y^2 = -OH$).

The pinanediol boronic acid ester and the pinacol boronic acid ester are abbreviated "-C10H16" and

"-C6H12" respectively. Other illustrations of diols useful for deriving boronic acid esters are 1,2-ethanediol, 1,3-propanediol, 1,2-propanediol, 2,3-butanediol, 1,2-diisopropylethanediol, 5,6-decanediol, 1,2-dicyclohexylethanediol.

Note that throughout the text when an alkyl substituent is mentioned, the normal alkyl structure is meant (e.g. butyl is n-butyl) unless otherwise specified. However, in the definition of radicals above (e.g. R³), both branched and straight chains are included in the scope of alkyl.

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It is understood that many of the compounds of the present invention contain one or more chiral centers and that these stereoisomers may possess distinct physical and biological properties. The present invention comprises all of the stereoisomers or mixtures thereof. If the pure enantiomers or diastereomers are desired, they may be prepared using starting materials with the appropriate stereochemistry, or may be separated from mixtures of undesired stereoisomers by standard techniques, including chiral chromatography and recrystallization of diastereomeric salts.

Synthesis

The compounds of formula (I) can be prepared using
the reactions and techniques described below. The
reactions are performed in a solvent appropriate to the
reagents and materials employed and suitable for the
transformations being affected. It will be understood
by those skilled in the art of organic synthesis that
the functionality present on the molecule should be
consistent with the chemical transformations proposed
and this will sometimes require judgment as to the order
of synthetic steps or selection of particular process
scheme used from that shown below in order to obtain a
desired compound of the invention.

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Scheme 1. Synthesis of Thrombin Inhibitors

Reagents: a. IBCF, NMM, RCO₂H, Et₃N, 0 °C, b. NaN₃, c. H₂, Pd(OH)₂/C, HCl, d. DMAP, aminoiminomethanesulfonic acid, e. phenylboronic acid

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Amine hydrochloride 1 is readily available via the procedure of Kettner and Shenvi (EP 0293881 A2).

There are numerous synthetic methods by which to prepare amide 2, however, competing with amide formation is the cyclization of 1 to afford a complex mixture containing the desired amide and the corresponding N-acylboroproline. Since purification at this stage is unfeasible, choosing the correct method for amide formation is crucial to obtaining 2 in a purity suitable for subsequent synthetic transformations.

Three methods are preferred for the preparation of 2. 10 In the first, a solution of 1 in tetrahydrofuran or dichloromethane at 0 °C is treated sequentially with the desired acid chloride followed by two equivalents of triethylamine. The mixture is then allowed to warm to room temperature overnight. The second method is the mixed anhydride procedure of Anderson, et. al. (J. Am. 15 Chem. Soc. 1967, 89, 5012). In this method the isobutyl mixed anhydride is generated by dissolving the carboxylic acid component in tetrahydrofuran and adding one equivalent of N-methylmorpholine. The solution is 20 cooled to 0 °C and one equivalent of isobutyl chloroformate is added. After 5 minutes, a solution of 1 in chloroform is added, followed by the addition of one equivalent of triethylamine. The mixture is typically stirred at 0 °C for one hour followed by one 25 to several hours at room temperature. The third method for amide formation is the hydroxybenzotriazole/DCC method of König and Geiger (Chem. Ber. 1970, 103, 788-Thus, to a solution of 1 and the carboxylic acid component in dimethylformamide or tetrahydrofuran at 0

The preferred method for the preparation of azide 3 is by reaction of 2 with sodium azide (1.1 eq) in dimethylformamide at 70 °C for 2 hours.

°C is added N-methylmorpholine, 1-hydroxybenzotriazole

hydrate (2 eq) and DCC (1.05 eq). The solution is

allowed to warm to room temperature overnight.

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The azide displacement may also be performed prior to amide formation. This is the preferred method in cases where the rate of amide formation is slow relative to the rate of cyclization. Azide 4 is prepared by a modification of the procedure of Kettner and Shenvi (EP 0293881 A2) as shown in Scheme 2. Thus, bromide 5 is reacted with sodium azide, followed by homologation to give 6, chloride displacement to afford 7 and acidic hydrolysis to give 6. Amide formation between 4 and the carboxylic acid component then affords 3 directly.

Scheme 2. Synthesis of Azide 4

Reagents: a. NaN₃ b. CHCl₂Li, ZnCl₂, c. LiN(TMS)₂, d. 4M HCl, dioxane

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Reduction of azide 3 to amine 8 may be accomplished by hydrogenation over precious metal catalysts. The preferred catalyst for this transformation is Pearlman's catalyst (palladium hydroxide on carbon). The amine is typically isolated as the hydrochloride salt. Isolation of 8 as the free base typically results in lowered yields. Salts of 8 which may confer superior physical properties may be preferred over the hydrochloride salt.

Formamidination of amine 8 may be accomplished using 25 cyanamide. Due to the low reactivity of amine 8,

however, the preferred method for this transformation is reaction with 4-dimethylaminopyridine (DMAP) and aminoiminomethanesulfonic acid (AMSA, prepared by the method of Kim, et. al., *Tetrahedron Lett.* 1988, 29, 3183-6). This affords guanidine 9, which is isolated as the bisulfite or hydrochloride salt.

Cleavage of pinanediol ester 9 may be accomplished using anhydrous boron trichloride according to the procedure of Matteson and Ray (J. Am. Chem. Soc. 1980, 102, 7588). This method, however, is strongly Lewis acidic and leads to partial destruction of the substrate. The preferred method for water soluble boronic acids is a transesterification reaction that is run in the presence of excess phenylboronic acid. The free boronic acid 10 may then be isolated using cation exchange chromatography.

The isothiouronium functionalized analogs 11/12 are prepared from bromide 2 according to the procedure of Kettner and Shenvi (EP 0293881 A2).

Inhibitors containing a sulfonamide in place of a carboxamide are prepared from either 1 or 4 by reaction with a sulfonyl chloride in the presence of a hindered amine (Scheme 3). The product sulfonamide 13 is then converted to the guanidinium 14 or isothiouronium 15 in the same manner as the corresponding carboxamides.

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Scheme 3. Synthesis of Sulfonamides

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Inhibitors containing the borolysine moiety are prepared analogously to those containing boroarginine according to Kettner and Shenvi (EP 0293881 A2).

Novel biaryls synthesized in this invention are prepared through palladium catalyzed coupling of an appropriate arylmetal species to the aryl halide of choice using the methods described in Negishi, et. al., Org. Synth. 1987, 66, 67-74, and references cited within.

20

EXAMPLE 1: N^{1} -(4-Phenylbenzoyl)boroarginine (+)-Pinanediol, Bisulfite

Part A: (+)-Pinanediol 4-bromo-1(R)-(4-phenylbenzoy1) aminobutane-1-boronate. To a solution of (+)pinanediol 4-bromo-1(R)-aminobutane-1-boronate hydrochloride (5.00 g, 13.6 mmol) in dichloromethane (50 mL) at 0 °C was added 4-phenylbenzoyl chloride (4.97 g, 22.9 mmol) followed by N-methylmorpholine (4 mL, 36 mmol). After 1 hour, the cooling bath was removed and the mixture stirred at room temperature for 2 hours. The mixture was then diluted with ethyl acetate and 10 washed with 0.1 M hydrochloric acid, saturated sodium bicarbonate and saturated sodium chloride. The organic phase was dried over anhydrous magnesium sulfate, filtered and the filtrate concentrated in vacuo to afford 3.37 g (48%) of the desired amide, mass spectrum: $(M+H)^+ = 510/512$; ¹H NMR (300 MHz, CDCl₃) $\delta 7.9$ (2H, d, J 15 = 8.3), 7.84 (1H, bs), 7.6 (2H, d, J = 8.3), 7.44 (5H, m), 4.37 (1H, m), 3.41 (1H, t, J = 6.9), 2.0 (10H, m) 1.49 (3H, s), 1.38 (1H, m), 1.29 (3H, s), 0.91 (3H, s).

20 Part B: (+)-Pinanediol 4-azido-1(R)-(4-phenylbenzoyl)aminobutane-1-boronate. To a solution of (+)pinanediol 4-bromo-1(R)-(4-phenylbenzoyl)aminobutane-1boronate (3.37 g, 6.60 mmol) in dimethylformamide (6 mL) was added sodium azide (547 mg, 8.41 mmol). resulting mixture was heated at 70 °C for 2 hours, cooled to room temperature, and diluted with ethyl The mixture was then washed with water, saturated sodium chloride and dried over anhydrous magnesium sulfate. Filtration, followed by 30 concentration of the filtrate in vacuo gave 3.04 g (97%) of the desired azide, mass spectrum: $(M+H)^+ = 473$; 1H NMR (300 MHz, CDCl₃) δ 7.89 (2H, d, J = 8.3), 7.75 (1H, bs), 7.3 (7H, m), 4.32 (1H, m), 3.32 (1H, m), 2.0 (10H, m) 1.48 (3H, s), 1.3 (4H, m), 0.9 (3H, s).

Part C: N^{1} – (4-Phenylbenzovl) boroornithine (+) – pinanediol, hydrochloride. To a solution of (+)pinanediol 4-azido-1(R)-(4-phenylbenzoyl)aminobutane-1boronate (3.04 g, 6.44 mmol) in methanol (30 mL) was added Pearlman's catalyst Pd(OH)2/C, 200 mg) and 1 M hydrochloric acid (6.5 mL, 6.5 mmol). The mixture was placed on a Parr apparatus and hydrogenated at 50 psi The mixture was filtered using Celite™, for 3 hours. washed with methanol and the filtrate concentrated in The resulting amorphous solid was dissolved in water and washed with ether. The aqueous phase was then concentrated in vacuo and crystallized from ethyl acetate-hexanes, giving 1.52 g (49%) of the desired amine hydrochloride, mass spectrum: $(M+H)^+ = 447$; mp: 157-170 °C; ¹H NMR (400 MHz, CDCl₃/DMSO-d₆) δ 9.88 (1H, bs), 8.18, (2H, d, J = 8.3), 8.13 (3H, bs), 7.68 (2H, d, bs)J = 8.3), 7.61 (2H, d J = 7.0), 7.45 (2H, d, J = 7.0), 7.37 (1H, d, J = 7.30), 4.20 (1H, d, J = 6.3), 2.99 (1H, m), 2.87 (2H, m), 2.31 (1H, m), 2.13 (1H, m), 1.84 (7H, m), 1.56 (1H, d, J = 10.0), 1.42 (3H, s), 1.29 (3H, s), 0.89 (3H, s).

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Part D: N^{1} -(4-Phenylbenzoyl)boroarginine (+)pinanediol, bisulfite. To a solution of N^{1} -(4-25 phenylbenzoyl)boroornithine (+)-pinanediol, hydrochloride (80 mg, 0.17 mmol) in ethanol (2 mL) was added 4-dimethylaminopyridine (40 mg, 0.33 mmol). After 15 minutes, aminoiminomethanesulfonic acid (40 mg, 0.32 mmol) was added and the resulting mixture heated at 30 reflux for 3 hours. After cooling to room temperature, the mixture was filtered and the filtrate concentrated The residue was dissolved in chloroform and washed with 0.1 M hydrochloric acid, water and dried over anhydrous magnesium sulfate. Filtration, followed 35 by concentration of the filtrate in vacuo afforded 73 mg

(84%) of the desired guanidine, mass spectrum: $(M+H)^+ = 489$; ¹H NMR (400 MHz, CDCl₃, 60 °C) δ 9.48 (1H, bs), 8.10 (2H, d, J = 8.1), 8.07 (1H, bs), 7.75 (1H, bs), 7.54 (2H, d, J = 8.3), 7.48 (2H, d, J = 7.0), 7.35 (3H, m), 7.06 (4H, bs), 4.19 (1H, bd, J = 8.3), 3.1 (2H, m), 2.84 (1H, m), 2.29 (1H, m), 2.12 (1H, m), 1.96 (1H, m), 1.75 (6H, m), 1.47 (1H, d, J = 10.2), 1.40 (3H, s), 1.24 (3H, s), 0.83 (3H, s).

10 EXAMPLE 34: (+)-Pinanediol 4-(Formamidino)thio-1(R)-(4-phenylbenzoyl)aminobutane-1-boronate, Hydrobromide

(+)-Pinanediol 4-(formamidino)thio-1(R)-(4phenylbenzoyl) aminobutane-1-boronate, hydrobromide. 15 a solution of (+)-pinanediol 4-bromo-1(R)-(4-phenylbenzoyl)aminobutane-1-boronate (200 mg, 0.392 mmol) in methanol (3 mL) was added thiourea (120 mg, 1.58 mmol). The reaction was stirred at room temperature for 3 days. The mixture was concentrated in vacuo, the residue 20 dissolved in water and washed with ether. Concentration of the aqueous portion afforded 80 mg (35%) of the desired isothiourea, mass spectrum: $(M+H)^+ = 506$; ¹H NMR (300 MHz, CDCl₃) $\delta 8.15$ (2H, d, J = 8.4), 7.61 (2H, d, J = 8.4), 7.52 (2H, m), 7.38 (3H, m), 6.47 (1H, bs), 4.2325 (1H, dd, J = 6.6, 1.9), 3.24 (1H, m), 3.14, (1H, m),2.96, (1H, m), 2.32 (1H, m), 2.15 (1H, m), 1.99 (1H, m), 1.78 (6H, m), 1.48 (1H, d, J = 10.1), 1.42 (3H, s), 1.27 (3H, s), 0.86 (3H, s).

30 The compounds listed in Tables 1-12 can be prepared using the above examples.

TABLE 1

where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is $X(CH_2)_3$ -, A

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| | Ex | . x | $\mathbf{R}^{\mathbf{A}}$ | $R^{\mathbf{B}}$ | RC | $\mathbf{Y}^{1},\mathbf{Y}^{2}$ | Phys |
|----|------|--------------------------|---------------------------|------------------|------------|---------------------------------|------|
| | | | | | | | Data |
| | 1 | NHC (NH) NH ₂ | н | н | . Ph | (+)-pinanediol | A |
| 10 | 2 | NHC (NH) NH ₂ | н | Ph | . Н | (+)-pinanediol | |
| | 3 | NHC (NH) NH2 | н | OPh | Ph | (+)-pinanediol | В |
| | 4 | NHC (NH) NH ₂ | н | н | 4-pyridyl | (+)-pinanediol | С |
| | · 5 | NHC (NH) NH2 | COPh | Н | H | (+)-pinanediol | |
| | 6 | NHC (NH) NH2 | н | COPh | н | (+)-pinanediol | |
| 15 | 7 | NHC (NH) NH2 | H | Н | COPh | (+)-pinanediol | |
| | 8 | NHC (NH) NH ₂ | н | NHCbz | н | (+)-pinanediol | |
| | 9 | NHC (NH) NH2 | н | NMeCbz | Н | (+)-pinanediol | |
| | 10 | NHC (NH) NH ₂ | н | Н | Et | (+)-pinanediol | |
| | 11 | NHC (NH) NH2 | . н | H | n-Pr | (+)-pinanediol | • |
| 20 | . 12 | NHC (NH) NH2 | н | н | i-Pr | (+)-pinanediol | |
| | 13 | NHC (NH) NH2 | н | H | n-Bu | (+)-pinanediol | • |
| | 14 | NHC (NH) NH2 | н | H | t-Bu | (+)-pinanediol | |
| | 15 | NHC (NH) NH2 | н | н | n-hexyl | (+)-pinanediol | |
| | 16 | NHC (NH) NH2 | Н | н | cyclohexyl | (+)-pinanediol | |
| 25 | 17 | NHC (NH) NH2 | NHCO (CH2) 2Ph | н | н | (+)-pinanediol | |

| | 18 | NHC (NH) NH2 | H | | O-n-Bu | (+)-pinanediol | |
|----|----|--------------------------|-----------------------------|--------|-------------------------------------|----------------|------|
| | 19 | NHC (NH) NH ₂ | Ĥ | | NHCOcyclopropyl | (+)-pinanediol | |
| | Ex | x | $R^{\mathbf{A}}$ | RB | RC | Y^1, Y^2 | Phys |
| | | | | | | | Data |
| 5 | | | • | | | | |
| | 20 | NHC (NH) NH2 | . Н | Н | NHCO-cyclohexyl | (+)-pinanediol | |
| | 21 | NHC (NH) NH2 | H | н | NHCO (4-C6H4OMe) | (+)-pinanediol | |
| | 22 | NHC (NH) NH2 | H | н | 4-C ₆ H ₄ OMe | (+)-pinanediol | |
| | 23 | NHC (NH) NH2 | CO_2CH_2 (2- C_6H_4Ph) | , Н | H | (+)-pinanediol | |
| 10 | 24 | NHC (NH) NH ₂ | н | н | 1-naphthyl | (+)-pinanediol | |
| | 25 | NHC (NH) NH ₂ | H | Н | 4-C6H4CO2H | (+)-pinanediol | |
| | 26 | NHC (NH) NH2 | COPh | н | Me | (+)-pinanediol | |
| | 27 | NHC (NH) NH2 | н | NHCbz | n-Bu | (+)-pinanediol | • |
| | 28 | NHC (NH) NH2 | н | NMeCbz | n-Bu | (+)-pinanediol | |
| 15 | 29 | NHC (NH) NH2 | Me | Н | Ph | (+)-pinanediol | QQ |
| | 30 | NHC (NH) NH2 | Me | . н | 4-C6H4CO2H | (+)-pinanediol | |
| | 31 | NHC (NH) NH2 | Н | н | 4-C6H4CO2Me | (+)-pinanediol | |
| | 32 | NHC (NH) NH2 | Me | н | 4-C6H4CO2Me | (+)-pinanediol | |
| • | 33 | NHC (NH) NH ₂ | Н | OMe | Ph | (+)-pinanediol | |
| 20 | 34 | SC (NH) NH2 | н | Н | Ph | (+)-pinanediol | D |
| | 35 | SC (NH) NH ₂ | н | Ph | Н | (+)-pinanediol | E |
| | 36 | SC (NH) NH2 | Н | OPh | Н | (+)-pinanediol | F |
| | 37 | SC (NH) NH ₂ | COPh | Н | Н | (+)-pinanediol | G |
| | 38 | SC (NH) NH2 | н | COPh | Н | (+)-pinanediol | H |
| 25 | 39 | SC (NH) NH ₂ | н | н | COPh | (+)-pinanediol | I |
| | 40 | SC (NH) NH ₂ | н | NHCbz | Н | (+)-pinanediol | J |
| | 41 | SC (NH) NH ₂ | н | NMeCbz | . Н | (+)-pinanediol | ĸ |
| | 42 | SC (NH) NH ₂ | Н | н | Et | (+)-pinanediol | r. |
| | 43 | SC (NH) NH ₂ | Н | Н | n-Pr | (+)-pinanediol | М |
| 30 | 44 | SC (NH) NH2 | . Н | H | i-Pr | (+)-pinanediol | N |
| | 45 | SC (NH) NH2 | н | H | n-Bu | (+)-pinanediol | 0 |
| | 46 | SC (NH) NH ₂ | н | н | t-Bu | (+)-pinanediol | P |
| | 47 | SC (NH) NH ₂ | н | н | n-hexyl | (+)-pinanediol | Q |
| | 48 | SC (NH) NH2 | н | H | cyclohexyl | (+)-pinanediol | R |
| 35 | 49 | SC (NH) NH2 | NHCOCH2CH2Ph | Н | H | (+)-pinanediol | s |
| | 50 | SC (NH) NH ₂ | H | Н | O-n-Bu | (+)-pinanediol | T · |
| | | | | | | • | |

| • | 51 | SC (NH) NH ₂ | н | H | NHCOcyclopropyl | (+)-pinanediol | U |
|----|----|---------------------------------|-----------------------------|--------|------------------|---------------------------------|------------|
| | Ex | x | RA | RB | RC | Y ¹ , Y ² | Phys |
| | | | , | | • | | Data |
| 5 | 52 | SC (NH) NH ₂ | Н | н | NHCOcyclohexyl | (+)-pinanediol | v |
| | 53 | SC (NH) NH ₂ | н | Н | NHCO (4-C6H4OMe) | (+)-pinanediol | W |
| | 54 | SC (NH) NH ₂ | н | H | 4-C6H4OMe | (+)-pinanediol | , X |
| | 55 | SC (NH) NH ₂ | CO_2CH_2 (2- C_6H_4Ph) | Н | Н | (+)-pinanediol | Y |
| | 56 | SC (NH) NH ₂ | н | Н | 1-naphthyl | (+)-pinanediol | |
| 10 | 57 | SC (NH) NH ₂ | н | Н | 4-C6H4CO2H | (+)-pinanediol | |
| | 58 | SC (NH) NH ₂ | н | NHCbz | n-Bu | (+)-pinanediol | Z |
| | 59 | SC (NH) NH ₂ | н | NMeCbz | n-Bu | (+)-pinanediol | AA |
| | 60 | SC (NH) NH ₂ | COPh | н | Me | (+)-pinanediol | BB |
| | 61 | SC (NH) NH ₂ | H | Н | 4-pyridyl | (+)-pinanediol | |
| 15 | 62 | SC (NH) NH ₂ | Me | Н | 4-C6H4CO2H | (+)-pinanediol | |
| | 63 | SC (NH) NH ₂ | н | H | 4-C6H4CO2Me | (+)-pinanediol | |
| | 64 | SC (NH) NH ₂ | Me | н | 4-C6H4CO2Me | (+)-pinanediol | |
| | 65 | SC (NH) NH2 | Me | H | Ph | (+)-pinanediol | |
| | 66 | SC (NH) NH ₂ | н | OMe | Ph | (+)-pinanediol | |
| 20 | 67 | CH2NH2 | H | н | Ph | (+)-pinanediol | |
| | 68 | CH2NH2 | H | Ph | н | (+)-pinanediol | |
| | 69 | CH2NH2 | н | OPh | H | (+)-pinanediol | |
| | 70 | CH2NH2 | COPh | H | H | (+)-pinanediol | |
| | 71 | CH2NH2 | н | COPh | H | (+)-pinanediol | |
| 25 | 72 | CH ₂ NH ₂ | н | н | COPh | (+)-pinanediol | |
| | 73 | CH2NH2 | · H | NHCbz | H | (+)-pinanediol | |
| | 74 | CH2NH2 | н | NMeCbz | н | (+)-pinanediol | |
| | 75 | CH2NH2 | . Н | H | Et | (+)-pinanediol | |
| | 76 | CH2NH2 | Н | н | n-Pr | (+)-pinanediol | |
| 30 | 77 | CH2NH2 | н | н | i-Pr | (+)-pinanediol | |
| | 78 | CH2NH2 | н | H | n-Bu | (+)-pinanediol | |
| | 79 | CH2NH2 | н | H | t-Bu | (+)-pinanediol | |
| | 80 | CH2NH2 | н | н | n-hexyl | (+)-pinanediol | |
| | 81 | CH2NH2 | н | Н | cyclohexyl | (+)-pinanediol | |
| 35 | 82 | CH2NH2 | NHCOCH2CH2Ph | Н | Н | (+)-pinanediol | |
| | 83 | CH2NH2 | н | . н | O-n-Bu | (+)-pinanediol | |

| | | | | | | • | |
|----|-----|--------------------------|--|------------|------------------|---------------------|------|
| | 84 | CH2NH2 | н | | NHCOcyclopropyl | (+)-pinanediol | |
| | Ex | х | $R^{\mathbf{A}}$ | RB | RC | Y ¹ , Y2 | Phys |
| | | | | | | | Data |
| | 85 | CH2NH2 | н | Н | NHCOcyclohexyl | (+)-pinanediol | |
| 5 | 86 | CH2NH2 | . Н | Н | NHCO (4-C6H4OMe) | (+)-pinanediol | |
| | 87 | CH2NH2 | Н | Н | 4-C6H4OMe | (+)-pinanediol | |
| | 88 | CH2NH2 | CO ₂ CH ₂ (2-C ₆ H ₄ Ph) | н | н | (+)-pinanediol | |
| | 89 | CH2NH2 | Н | Н | 1-naphthyl | (+)-pinanediol | |
| | 90 | CH2NH2 | H | н | 4-C6H4CO2H | (+)-pinanediol | |
| 10 | 91 | CH2NH2 | H | NHCbz | n-Bu | (+)-pinanediol | |
| | 92 | CH2NH2 | Н | NMeCbz | n-Bu | (+)-pinanediol | |
| | 93 | CH2NH2 | COPh | н | Me | (+)-pinanediol | |
| | 94 | CH2NH2 | н | Н | 4-pyridyl | (+)-pinanediol | |
| | 95 | CH2NH2 | Me | н | 4-C6H4CO2H | (+)-pinanediol | |
| 15 | 96 | CH2NH2 | н | н | 4-C6H4CO2Me | (+)-pinanediol | |
| | 97 | CH2NH2 | Me | н | 4-C6H4CO2Me | (+)-pinanediol | |
| | 98 | CH2NH2 | Me | . H | Ph | (+)-pinanediol | |
| | 99 | CH2NH2 | н | OMe | Ph | (+)-pinanediol | |
| | 100 | CH2NH2 | н | OMe | Ph | н, н | |
| 20 | 101 | NHC (NH) NH2 | н | н | Ph | н, н | • |
| | 102 | NHC (NH) NH2 | н | Ph | Н | н, н | |
| | 103 | NHC (NH) NH2 | . н | · OPh | . Ph | н, н | |
| | 104 | NHC (NH) NH2 | ,H | н | 4-pyridyl | н, н | |
| | 105 | NHC (NH) NH2 | COPh | . н | н | н, н | |
| 25 | 106 | NHC (NH) NH2 | н | COPh | н | н, н | |
| | 107 | NHC (NH) NH2 | . н | H | COPh | н, н | |
| | 108 | NHC (NH) NH2 | H | NHCbz | H | н, н | |
| | 109 | NHC (NH) NH2 | н | NMeCbz | H | н, н | • |
| * | 110 | NHC (NH) NH2 | н | н | Et | Н, Н | • |
| 30 | 111 | NHC (NH) NH ₂ | . н | н | n-Pr | н, н | |
| • | 112 | NHC (NH) NH ₂ | н | . н | i-Pr | н, н | |
| | 113 | NHC (NH) NH2 | н | н | n-Bu | н, н | |
| • | 114 | NHC (NH) NH ₂ | н | H | t-Bu | н, н | • |
| | 115 | NHC (NH) NH ₂ | н | Н | n-hexyl | н, н | |
| 35 | 116 | NHC (NH) NH2 | н | н | cyclohexyl | н, н | • |
| | 117 | NHC (NH) NH ₂ | NHCO (CH2) 2Ph | H | н | н, н | |

| | Ex | x | RA | RB | RC | Y ¹ ,Y ² | Phys |
|----|-----|--------------|-------------------|--------|------------------|--------------------------------|--------|
| | | • | | | • | | Data . |
| | 118 | NHC (NH) NH2 | | н | 0-n-Bu | н, н | |
| | 119 | NHC (NH) NH2 | н | H | NHCOcyclopropyl | н, н | |
| 5 | 120 | NHC (NH) NH2 | н | н | NHCO-cyclohexyl | н, н | |
| | 121 | NHC (NH) NH2 | н | Н | NHCO (4-C6H4OMe) | н, н | |
| | 122 | NHC (NH) NH2 | н | н | 4-C6H4OMe | н, н | |
| • | 123 | NHC (NH) NH2 | CO2CH2 (2-C6H4Ph) | Н | H | Н, Н | |
| | 124 | NHC (NH) NH2 | н | H | 1-naphthyl | н, н | |
| 10 | 125 | NHC (NH) NH2 | н | H | 4-C6H4CO2H | н, н | |
| | 126 | NHC (NH) NH2 | COPh | . Н | Me | , н,н | |
| | 127 | NHC (NH) NH2 | н | NHCbz | n-Bu | н, н | |
| | 128 | NHC (NH) NH2 | н | NMeCbz | n-Bu | н, н | |
| | 129 | NHC (NH) NH2 | Me | H | Ph | н, н | |
| 15 | 130 | NHC (NH) NH2 | Me | Н | 4-C6H4CO2H | H, H | |
| | 131 | NHC (NH) NH2 | н | H | 4-C6H4CO2Me | н, н | |
| | 132 | NHC (NH) NH2 | Me | н | 4-C6H4CO2Me | Н, Н | |
| | 133 | NHC (NH) NH2 | н | OMe | Ph | Н, Н | |
| | 134 | SC (NH) NH2 | н | н | Ph | н, н | |
| 20 | 135 | SC (NH) NH2 | н | Ph | н | н, н | |
| | 136 | SC (NH) NH2 | н | OPh | н | Н, Н | . • |
| | 137 | SC (NH) NH2 | COPh | Н | н | н, н | |
| | 138 | SC (NH) NH2 | . н | COPh | н | н, н | • |
| | 139 | SC (NH) NH2 | н | H | COPh | H, H | |
| 25 | 140 | SC (NH) NH2 | н | NHCbz | · H | н, н | |
| | 141 | SC (NH) NH2 | н : | NMeCbz | н | н, н | |
| | 142 | SC (NH) NH2 | н | Н | Et | H, H | |
| | 143 | SC (NH) NH2 | н | Н | n-Pr | н,н | |
| | 144 | SC (NH) NH2 | н | H | i-Pr | Н, Н | |
| 30 | 145 | SC (NH) NH2 | . Н | H | n-Bu | . н, н | |
| | 146 | SC (NH) NH2 | H | Н | t-Bu | н, н | |
| | 147 | SC (NH) NH2 | Н | Н | n-hexyl | н, н | |
| | 148 | SC (NH) NH2 | Н | Н | cyclohexyl | . н, н | |
| | 149 | SC (NH) NH2 | NHCOCH2CH2Ph | н | H | н, н | |
| 35 | 150 | SC (NH) NH2 | н | н | O-n-Bu | н, н | • |
| | Ex | · x | RA | RB | _R C | Y^1, Y^2 | Phys |

| | | | | | | | Data |
|----|-----|-------------------------|-------------------|--------|---------------------------------|---------------------------------|------|
| | 151 | SC (NH) NH2 | н | н | NHCO (CH ₂) 2phenyl | Н, Н | RR |
| | 152 | SC (NH) NH ₂ | H | Н | NHCOcyclohexyl | Н, Н | |
| | 153 | SC (NH) NH2 | н | Н | NHCO (4-C6H4OMe) | Н, Н | |
| 5 | 154 | SC (NH) NH2 | н | Н | 4-C6H4OMe | н, н | |
| | 155 | SC (NH) NH ₂ | CO2CH2 (2-C6H4Ph) | Н | н | Н, Н | |
| | 156 | SC (NH) NH2 | H | Н | 1-naphthyl | Н, Н | |
| | 157 | SC (NH) NH2 | н | Н | 4-C6H4CO2H | Н, Н | |
| | 158 | SC (NH) NH ₂ | н | NHCbz | n-Bu | н, н | |
| 10 | 159 | SC (NH) NH ₂ | H | NMeCbz | n-Bu | Н, Н | |
| | 160 | SC (NH) NH2 | COPh | н | · Me | н, н | |
| | 161 | SC (NH) NH2 | н | Н | 4-pyridyl | Н, Н | |
| | 162 | SC (NH) NH ₂ | Me | н | 4-C6H4CO2H | . Н, н | |
| | 163 | SC (NH) NH2 | н | Н | 4-C6H4CO2Me | Н, Н | |
| 15 | 164 | SC (NH) NH2 | Me | Н | 4-C6H4CO2Me | н, н | |
| | 165 | SC (NH) NH2 | Me | н | Ph | н, н | |
| | 166 | SC (NH) NH2 | н | OMe | Ph | Н, Н | |
| • | 167 | CH2NH2 | н | Н | Ph | н, н | |
| | 168 | CH2NH2 | . н | Ph | H | . Н, Н | |
| 20 | 169 | CH2NH2 | н | OPh | н | Н, Н | |
| | 170 | CH2NH2 | COPh | Н | . Н | н, н | |
| | 171 | CH2NH2 | н | COPh | H | Н, Н | |
| | 172 | CH2NH2 | н | H | COPh | н, н | |
| | 173 | CH2NH2 | н | NHCbz | H · | Н,Н | |
| 25 | 174 | CH2NH2 | н | NMeCbz | Н | . н, н | |
| | 175 | CH2NH2 | , H | Н | Et | н, н | |
| | 176 | CH2NH2 | н | H | n-Pr | н, н | |
| | 177 | CH2NH2 | . н | H | i-Pr | н, н | |
| | 178 | CH2NH2 | н | H | n-Bu | н, н | |
| 30 | 179 | CH2NH2 | н | H | t-Bu | н, н | |
| | 180 | CH2NH2 | Н | | n-hexyl | н, н | |
| | 181 | CH2NH2 | Н | | cyclohexyl | Н, Н | |
| | 182 | CH2NH2 | NHCOCH2CH2Ph | | Н | Н, Н | |
| | 183 | CH2NH2 | Н | | O-n-Bu | н, н | |
| 35 | Ex | x | RA | RB | · _R C | Y ¹ , Y ² | Phys |
| | | | | | | | Data |

| Н, Н | NHCOcyclopropyl | Н | H | CH2NH2 | 184 | |
|------|------------------|--------|-----------------------------|--------|-----|----|
| н, н | NHCOcyclohexyl | н | н | CH2NH2 | 185 | |
| н, н | NHCO (4-C6H4OMe) | Н | . н | CH2NH2 | 186 | |
| н, н | 4-C6H4OMe | Н | H | CH2NH2 | 187 | |
| н, н | н | н | CO_2CH_2 (2- C_6H_4Ph) | CH2NH2 | 188 | 5 |
| н, н | 1-naphthyl | Н | н | CH2NH2 | 189 | • |
| н, н | 4-C6H4CO2H | Н | н | CH2NH2 | 190 | |
| н, н | n-Bu | NHCbz | н | CH2NH2 | 191 | |
| н, н | n-Bu | NMeCbz | н | CH2NH2 | 192 | |
| н, н | Ме | Н | COPh | CH2NH2 | 193 | 10 |
| H, H | 4-pyridyl | н | н | CH2NH2 | 194 | |
| H,H | 4-C6H4CO2H | Н | · Me | CH2NH2 | 195 | |
| н, н | 4-C6H4CO2Me | Н | н | CH2NH2 | 196 | |
| Н,Н | 4-C6H4CO2Me | Н | Me | CH2NH2 | 197 | |
| Н, Н | Ph | Н | Me | CH2NH2 | 198 | 15 |
| | | | | | | |

TABLE 2

where
$$R_1^2$$
 is $X(CH_2)_3$ -, and where R_1^3 is $X(CH_2)_3$ -, and $X(CH_2)_3$ -,

20

(

| | Ex | x | Y | Y^1, Y^2 | Phys |
|----|-----|--------------------------|--------|--------------------------------|------|
| | | | | | Data |
| | 199 | CH2NH2 | CO | (+)-pinanediol | |
| 25 | 200 | CH2NH2 | so_2 | (+)-pinanediol | |
| | 201 | NHC (NH) NH ₂ | СО | (+)-pinanediol | |
| ٠. | Ex | x | Y | Y ¹ ,Y ² | Phys |

| | | | | | Data |
|----|-----|--------------------------|-----------------|----------------|------|
| | 202 | NHC (NH) NH ₂ | so ₂ | (+)-pinanediol | |
| | 203 | SC (NH) NH2 | CO | (+)-pinanediol | cc |
| | 204 | SC (NH) NH ₂ | so ₂ | (+)-pinanediol | DD |
| 5 | 205 | CH2NH2 | СО | н, н | |
| | 206 | CH2NH2 | so ₂ | н, н | |
| | 207 | NHC (NH) NH ₂ | СО | Н, Н | |
| • | 208 | NHC (NH) NH ₂ | so ₂ | н, н | |
| | 209 | SC (NH) NH ₂ | со | н, н | |
| 10 | 210 | SC (NH) NH ₂ | so ₂ | н, н | |

TABLE 3

6

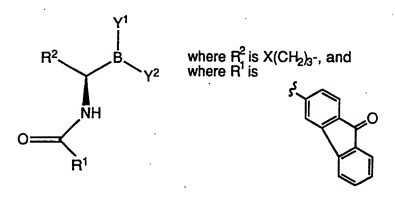
where
$$R^2$$
 is $XCH_2(CH_2)CH_2$, and where R^1 is

| 15 | | | | · | |
|----|-----|--------------------------|---|---------------------------------|------|
| | Ex | x | t | Y ¹ , Y ² | Phys |
| | | | | | Data |
| | 211 | NH ₂ | 2 | (+)-pinanediol | |
| • | 212 | SC (NH) NH2 | 2 | (+)-pinanediol | EE |
| 20 | 213 | SC (NH) NH2 | 1 | (+)-pinanediol | FF |
| | 214 | NHC (NH) NH2 | 2 | (+)-pinanediol | |
| | 215 | NHC (NH) NH ₂ | 1 | (+)-pinanediol | |
| | 216 | NH ₂ | 2 | н, н | |
| | 217 | SC (NH) NH ₂ | 2 | н, н | |
| 25 | | | | | |
| | Ex | x | T | ¥1,¥2 | Phys |

Data

| | 218 | SC (NH) NH ₂ | 1 . | н, н |
|---|-----|-------------------------|-----|------|
| | 219 | NHC (NH) NH2 | 2 | Н, Н |
| 5 | 220 | NHC (NH) NH2 | 1 | н, н |

TABLE 4



| 10 | Ex | x | Y^1, Y^2 | Phys Dat | ta |
|----|-----|-------------------------|----------------|----------|----|
| | 221 | CH2NH2 | (+)-pinanediol | | |
| | 222 | NHC (NH) NH2 | (+)-pinanediol | | |
| | 223 | SC (NH) NH ₂ | (+)-pinanediol | GG | |
| | 224 | CH2NH2 | Н, Н | | |
| 15 | 225 | NHC (NH) NH2 | н, н | | |
| | 226 | SC (NH) NH2 | H, H | | |

TABLE 5

where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is $X(R^2)_3$ -, and R^1

| | Ex | · x | Y^1, Y^2 | Phys Data |
|----|-----|-------------------------|----------------|-----------|
| 5 | 227 | CH2NH2 | (+)-pinanediol | |
| | 228 | NHC (NH) NH2 | (+)-pinanediol | |
| | 229 | SC (NH) NH ₂ | (+)-pinanediol | HH. |
| | 230 | CH2NH2 | н, н | |
| | 231 | NHC (NH) NH2 | Н, Н | • |
| 10 | 232 | SC (NH) NH2 | н, н | |

TABLE 6

where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is R^D
 R^1

15

()

| | Ex | x . | $R^{\mathbf{A}}$ | _R c | R^D | y ¹ , y ² | Phys | Data |
|-----|-----|--------------------------|---------------------------|----------------|-------------------|---------------------------------|------|------|
| | 233 | NHC (NH) NH2 | Me | Ph | OMe | (+)-pinanediol | | |
| | 234 | NHC (NH) NH2 | Me | Ph | CONH ₂ | (+)-pinanediol | | |
| | 235 | NHC (NH) NH2 | Me | Ph | F | (+)-pinanediol | | |
| . 5 | 236 | NHC (NH) NH ₂ | Me | Ph | CF3 | (+)-pinanediol | | |
| | 237 | NHC (NH) NH2 | Me | Ph | Cl | (+)-pinanediol | • | |
| | 238 | NHC (NH) NH2 | Me | Ph | . ОН | (+)-pinanediol | | |
| | 239 | NHC (NH) NH2 | Me | 4-C6H4CO2H | OMe | (+)-pinanediol | | |
| | 240 | NHC (NH) NH2 | Me | 4-С6Н4СО2Н | CONH2 | (+)-pinanediol | | |
| 10 | 241 | NHC (NH) NH2 | Me | 4-C6H4CO2H | F | (+)-pinanediol | | |
| | 242 | NHC (NH) NH2 | Me | 4-C6H4CO2H | CF3 | (+)-pinanediol | | |
| | 243 | NHC (NH) NH2 | Me | 4-C6H4CO2H | Cl | (+)-pinanediol | | |
| | 244 | NHC (NH) NH2 | Me | 4-C6H4CO2H | ОН | (+)-pinanediol | | |
| | 245 | SC (NH) NH2 | Me | Ph | OMe | (+)-pinanediol | | |
| 15 | 246 | SC (NH) NH2 | Me | Ph | CONH ₂ | (+)-pinanediol | | |
| | 247 | SC (NH) NH2 | Me | Ph | F | (+)-pinanediol | | |
| | 248 | SC (NH) NH2 | Me | Ph | CF3 | (+)-pinanediol | | |
| | 249 | SC (NH) NH ₂ | Me | Ph | Cl | (+)-pinanediol | | |
| | 250 | SC (NH) NH2 | Me | Ph | OH | (+)-pinanediol | | |
| 20 | 251 | SC (NH) NH ₂ | Me | 4-C6H4CO2H | OMe | (+)-pinanediol | | |
| | 252 | SC (NH) NH2 | Me | 4-С6Н4СО2Н | CONH ₂ | (+)-pinanediol | | |
| | 253 | SC (NH) NH ₂ | Me | 4-C6H4CO2H | F | (+)-pinanediol | | |
| | 254 | SC (NH) NH ₂ | Me | 4-C6H4CO2H | CF3 | (+)-pinanediol | | • |
| | 255 | SC (NH) NH ₂ | Me | 4-С644С02Н | Cl | (+)-pinanediol | • | |
| 25 | 256 | SC (NH) NH ₂ | Me | 4-C6H4CO2H | ОН | (+)-pinanediol | | |
| | 257 | CH2NH2 | Me | Ph | OMe | (+)-pinanediol | | |
| | 258 | CH2NH2 | Me | · Ph | CONH ₂ | (+)-pinanediol | | |
| | 259 | CH2NH2 | Me | Ph | F | (+)-pinanediol | | |
| | 260 | CH2NH2 | Me | Ph | CF3 | (+)-pinanediol | | |
| 30 | 261 | CH2NH2 | Me | Ph | Cl | (+)-pinanediol | | |
| | 262 | CH2NH2 | Me | Ph | ОН | (+)-pinanediol | | |
| | 263 | CH2NH2 | Me | 4-C6H4CO2H | OMe | (+)-pinanediol | | • |
| | 264 | CH2NH2 | Me | 4-C6H4CO2H | CONH2 | (+)-pinanediol | | |
| | 265 | CH2NH2 | Me | 4-C6H4CO2H | F | (+)-pinanediol | | |
| 35 | 266 | CH2NH2 | Me | 4-C6H4CO2H | CF3 | (+)-pinanediol | | |
| | Ex | x | $\mathbb{R}^{\mathbb{A}}$ | RC | ŔĎ | Y^1, Y^2 | Phys | Data |

| | 267 | CH2NH2 | Me | 4-C6H4CO2H | Cl | (+)-pinanediol | | |
|----|-----|---------------------------------|------|------------|-------------------|---------------------------------|-----------|--|
| | 268 | CH2NH2 | Ме | 4-С6Н4СО2Н | ОН | (+)-pinanediol | | |
| | 269 | NHC (NH) NH ₂ | Me | Ph | OMe | н, н | | |
| | 270 | NHC (NH) NH2 | Me | Ph | CONH ₂ | Н, Н | | |
| 5 | 271 | NHC (NH) NH2 | Me | Ph | F | Н, Н | | |
| | 272 | NHC (NH) NH2 | Me | Ph | CF3 | н, н | • | |
| | 273 | NHC (NH) NH2 | Me | Ph | Cl | н, н | | |
| | 274 | NHC (NH) NH2 | Me | Ph | ОН | Н, Н | | |
| | 275 | NHC (NH) NH2 | Me | 4-C6H4CO2H | OMe | н, н | • | |
| 10 | 276 | NHC (NH) NH2 | Me | 4-C6H4CO2H | CONH2 | н, н | | |
| | 277 | NHC (NH) NH2 | Me | 4-C6H4CO2H | F | н, н | | |
| | 278 | NHC (NH) NH2 | Me | 4-С6Н4СО2Н | CF3 | н, н | | |
| | 279 | NHC (NH) NH2 | Me | 4-C6H4CO2H | Cl | н, н | | |
| | 280 | NHC (NH) NH2 | Me | 4-C6H4CO2H | ОН | Н, Н | | |
| 15 | 281 | SC (NH) NH2 | Me | Ph | OMe . | н, н | | |
| | 282 | SC (NH) NH2 | Me | Ph | conh ₂ | н, н | | |
| | 283 | SC (NH) NH2 | Me | Ph | F | Н, Н | | |
| | 284 | SC (NH) NH ₂ | Me | Ph | CF3 | Н, Н | | |
| | 285 | SC (NH) NH ₂ | Me | Ph | Cl | н, н | | |
| 20 | 286 | SC (NH) NH ₂ | Me | Ph | OH | н, н | | |
| | 287 | SC (NH) NH ₂ | Me | 4-С6Н4СО2Н | OMe | н, н | | |
| | 288 | SC (NH) NH ₂ | Me | 4-С6Н4СО2Н | CONH ₂ | н, н | | |
| | 289 | SC (NH) NH ₂ | Me | 4-С6Н4СО2Н | F | н, н | | |
| | 290 | SC (NH) NH ₂ | . Me | 4-С6Н4СО2Н | CF3 | н, н | • | |
| 25 | 291 | SC (NH) NH ₂ | Me | 4-С6Н4СО2Н | Cl | н, н | | |
| | 292 | SC (NH) NH2 | Me | 4-C6H4CO2H | OH | н, н | | |
| | 293 | CH ₂ NH ₂ | Me | Ph | OMe | н, н | | |
| | 294 | CH ₂ NH ₂ | Me | Ph | conh ₂ | н, н | | |
| | 295 | CH ₂ NH ₂ | Me | Ph | F | н, н | | |
| 30 | 296 | CH ₂ NH ₂ | Me | Ph | CF3 | н, н | | |
| | 297 | CH2NH2 | Me | Ph | Cl | н, н | • | |
| | 298 | CH2NH2 | Me | Ph | ОН | н, н | | |
| | 299 | CH2NH2 | Me | 4-C6H4CO2H | OMe | Н, Н | | |
| | 300 | CH2NH2 | Me | 4-C6H4CO2H | CONH2 | н, н | | |
| 35 | En | ж | RA | RC | RD | Y ¹ , Y ² | Phys Data | |
| | 301 | CH2NH2 | Me | 4-C6H4CO2H | F | н, н | | |

| 302 | CH2NH2 | Me | 4-C6H4CO2H | CF3 | н, н |
|-----|--------|----|------------|-----|------|
| 303 | CH2NH2 | Me | 4-C6H4CO2H | Cl | н, н |
| 304 | CH2NH2 | Me | 4-C6H4CO2H | ОН | н, н |

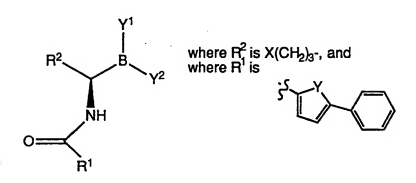
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TABLE 7

| | Ex | . X | $\mathbf{Y}^{1},\mathbf{Y}^{2}$ | Phys Data |
|----|-----|-------------------------|---------------------------------|-----------|
| | 305 | NHC (NH) NH2 | (+)-pinanediol | |
| 10 | 306 | SC (NH) NH ₂ | (+)-pinanediol | II |
| | 307 | CH2NH2 | (+)-pinanediol | · |
| | 308 | NHC (NH) NH2 | н, н | |
| | 309 | SC (NH) NH2 | н,н | |
| | 310 | CH2NH2 | Н,Н | |
| | | | | |

15

TABLE 8



| Ex | x | ¥ | Y^1, Y^2 | Phys | Data |
|-----|---|---|--|---|---|
| 311 | NHC (NH) NH2 | 0 | (+)-pinanediol | | |
| 312 | SC (NH) NH ₂ | 0 | (+)-pinanediol | | JJ |
| 313 | CH2NH2 | 0 | (+)-pinanediol | | |
| 314 | NHC (NH) NH2 | s | (+)-pinanediol | | |
| 315 | SC (NH) NH2 | s | (+)-pinanediol | | |
| 316 | CH2NH2 | S | (+)-pinanediol | | |
| 317 | NHC (NH) NH2 | 0 | н, н | | |
| 318 | SC (NH) NH ₂ | 0 | н, н | | |
| 319 | CH2NH2 | 0 | н, н | | |
| 320 | NHC (NH) NH2 | S | н, н | | |
| 321 | SC (NH) NH ₂ | s | н, н | | |
| 322 | CH2NH2 | S | н, н | | |
| | 311 312 313 314 315 316 317 318 319 320 321 | 311 NHC (NH) NH2 312 SC (NH) NH2 313 CH2NH2 314 NHC (NH) NH2 315 SC (NH) NH2 316 CH2NH2 317 NHC (NH) NH2 318 SC (NH) NH2 319 CH2NH2 319 CH2NH2 320 NHC (NH) NH2 321 SC (NH) NH2 | 311 NHC (NH) NH2 O 312 SC (NH) NH2 O 313 CH2NH2 O 314 NHC (NH) NH2 S 315 SC (NH) NH2 S 316 CH2NH2 S 317 NHC (NH) NH2 O 318 SC (NH) NH2 O 319 CH2NH2 O 320 NHC (NH) NH2 S 321 SC (NH) NH2 S | 311 NHC (NH) NH2 O (+) -pinanediol 312 SC (NH) NH2 O (+) -pinanediol 313 CH2NH2 O (+) -pinanediol 314 NHC (NH) NH2 S (+) -pinanediol 315 SC (NH) NH2 S (+) -pinanediol 316 CH2NH2 S (+) -pinanediol 317 NHC (NH) NH2 O H, H 318 SC (NH) NH2 O H, H 319 CH2NH2 O H, H 320 NHC (NH) NH2 S H, H 321 SC (NH) NH2 S H, H | 311 NHC (NH) NH2 O (+) -pinanediol 312 SC (NH) NH2 O (+) -pinanediol 313 CH2NH2 O (+) -pinanediol 314 NHC (NH) NH2 S (+) -pinanediol 315 SC (NH) NH2 S (+) -pinanediol 316 CH2NH2 S (+) -pinanediol 317 NHC (NH) NH2 O H, H 318 SC (NH) NH2 O H, H 319 CH2NH2 O H, H 320 NHC (NH) NH2 S H, H 321 SC (NH) NH2 S H, H |

TABLE 9

15

where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is $X(CH_2)_3$ -, and $X(CH_2)_3$ -, and

 R^B RC Y^1, Y^2 Ex X Phys Data 323 NHC (NH) NH2 H Ph (+)-pinanediol 324 NHC (NH) NH2 OBn H (+)-pinanediol · 20 325 SC (NH) NH2 Ph H (+)-pinanediol KK 326 SC (NH) NH2 H OBn (+)-pinanediol LL 327 CH2NH2 Н Ph (+)-pinanediol 328 CH2NH2 OBn Н (+)-pinanediol 329 NHC (NH) NH2 Н Ph H, H 25 330 NHC (NH) NH2 OBn Н H,H 331 SC (NH) NH2 H Ph H, H RC Y^1, Y^2 $R^{\mathbf{B}}$ Ex Phys Data 332 SC (NH) NH2 Н OBn H,H

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| 333 | CH2NH2 | H | Ph | н, н |
|-----|--------|-----|----|------|
| 334 | CH2NH2 | OBn | Н | н,н |

TABLE 10

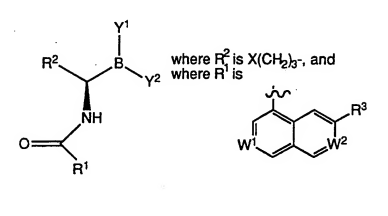
where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is $X(CH_2)_3$ -, and $X(CH_2)_3$ -, and

Y1, Y2 Ex X Phys Data 335 NHC (NH) NH2 (+)-pinanediol 336 SC (NH) NH2 (+)-pinanediol MM 10 337 (+)-pinanediol CH2NH2 338 NHC (NH) NH2 H,H 339 SC (NH) NH2 H,H 340 CH2NH2 H,H

15

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TABLE 11



Ex $X W^1 W^2 R^3 Y^1, Y^2$ Phys

| Data | | | | | | | |
|------|----------------|------|----|----|---------------------------------|-----|----|
| | (+)-pinanediol | Н | СН | N | NHC (NH) NH2 | 341 | |
| | (+)-pinanediol | Н | СН | N | SC (NH) NH ₂ | 342 | |
| , | (+)-pinanediol | Н | CH | N | CH ₂ NH ₂ | 343 | |
| | (+)-pinanediol | Ph | N | СН | NHC (NH) NH2 | 344 | 5 |
| 00 | (+)-pinanediol | Ph | N | СН | SC (NH) NH2 | 345 | |
| | (+)-pinanediol | Ph - | N | CH | CH2NH2 | 346 | |
| | H, H | Н | СН | N | NHC (NH) NH2 | 347 | |
| | н, н | H | CH | N | SC (NH) NH ₂ | 348 | |
| | н, н | Н | СН | N | CH ₂ NH ₂ | 349 | 10 |
| | н, н | Ph | N | CH | NHC (NH) NH2 | 350 | |
| | н, н | Ph | N | СН | SC (NH) NH2 | 351 | |
| | н.н | Ph | N | CH | CH2NH2 | 352 | |

TABLE 12

| R^2 B Y^2 | where R^2 is $X(CH_2)_3$ -, and where R^1 is |
|------------------|--|
| O R ¹ | N N N N N N N N N N N N N N N N N N N |

| Data | Phys | Y ¹ , Y ² | x | Ex | |
|------|------|---------------------------------|---------------------------------|--------|----|
| | • | (+)-pinanediol | NHC (NH) NH2 | 353 | |
| PP | | (+)-pinanediol | SC (NH) NH2 | 354 | |
| | | (+)-pinanediol | CH ₂ NH ₂ | 20 355 | 20 |
| | | н, н | NHC (NH) NH2 | 356 | |
| | | . н,н | SC (NH) NH2 | 357 | |
| | | н, н | CH2NH2 | 358 | |
| | | | | | |

15

WO 94/21650

TABLE 13

where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is

Ex X R³ Y¹, Y² Phys Data
359 SC(NH)NH2 H (+)-pinanediol NN

5

TABLE 14

where
$$R^2$$
 is $X(CH_2)_3$ -, and where R^1 is $(CH_2)_m$
 R^1

Where R^2 is $X(CH_2)_3$ -, and $(CH_2)_m$
 R^A

 Y^1, Y^2 10 $\mathbb{R}^{\mathbb{A}}$ R^B $\mathbb{R}^{\mathbb{C}}$ Ex Phys Data SC (NH) NH2 Н NHCO (CH2) 2Ph Н (+)-pinanediol RR SC (NH) NH2 Ph Н Н (+)-pinanediol SC (NH) NH2 2 Н OPh Ph (+)-pinanediol SC (NH) NH2 1 Н H 4-pyridyl (+)-pinanediol 15 NHC (NH) NH2 1 COPh H (+)-pinanediol NHC (NH) NH2 3 H COPh Н (+)-pinanediol NHC (NH) NH2 3 Н Н COPh (+),-pinanediol

Physical Data for Tables 1-14
A: MS (M+H) + = 489; ¹H NMR (400 MHz, CDCl₃, 60 °C)
9.48 (1H, bs), 8.10 (2 H, d, J = 8.1), 8.07 (1 H,
bs), 7.75 (1 H, bs), 7.54 (2 H, d, J = 8.3), 7.48 (2 H, d, J = 7.0), 7.35 (3 H, m), 7.06 (4 H, bs), 4.19
(1 H, bd, J = 8.3), 3.1 (2 H, m), 2.84 (1 H, m), 2.29
(1 H, m), 2.12 (1 H, m), 1.96 (1 H, m), 1.75 (6 H, m), 1.47 (1 H, d, J = 10.2), 1.40 (3 H, s), 1.24 (3
10 H, s), 0.83 (3 H, s).

B: MS (DCI - NH₃), 505 $(M + H)^+$.

C: MS $(M+H)^+ = 490$.

D: MS (M+H)⁺ = 506; ¹H NMR (300 MHz, CDCl₃) 8.15 (2 H, d, J = 8.4), 7.61 (2 H, d, J = 8.4), 7.52 (2 H, m), 7.38 (3 H, m), 6.47 (1 H, bs), 4.23 (1 H, dd, J = 6.6, 1.9), 3.24 (1 H, m), 3.14, (1 H, m), 2.96, (1 H, m), 2.32 (1 H, m), 2.15 (1 H, m), 1.99 (1 H, m), 1.78 (6 H, m), 1.48 (1 H, d, J = 10.1), 1.42 (3 H, s), 1.27 (3 H, s), 0.86 (3 H, s).

E: mp 145-150 °C.

25 F: MS (DCI - NH₃), 522 (M + H)⁺.

G: HRMS (DCI - NH₃), Calc: 534.2597, Found: 534.2609.

30 H: HRMS (DCI - NH₃), Calc: 534.2597, Found: 534.2605.

I: HRMS (DCI - NH₃), Calc: 534.2597, Found: 534.2609.

J: $[a]_D = -14.85^\circ$ (c = 0.606, MeOH); ¹H NMR (300 MHz, 35 DMSO - d₆) 10.07 (br s, 1 H), 10.05 (br s, 1 H), 8.96 (4 H, br s), 8.08 (1 H, s), 7.71 (1 H, dd, J = 8.1,

1.1), 7.61 (1 H, d, J = 7.7), 7.30 - 7.50 (6 H, m), 5.18 (2 H, s), 4.08 (1 H, br d), 3.08 - 3.25 (2 H, m), 2.50 - 2.65 (1 H, m), 2.15 - 2.30 (1 H, m), 1.97 - 2.10 (1 H, m), 1.40 - 1.90 (8 H, m), 1.31 (3 H, s), 1.24 (3 H, s), 0.84 (3 H, s); IR (KBr) 2500 - 3700 (br), 1734, 1646, 1578, 1550, 1440, 1222, 1058 cm⁻¹; MS (CI - NH₃), m/e (%) 537.2 (10.2, M + H - H₂NCN)+), 429.0 (42.8), 277.0 (100); Anal. Calcd for $C_{30}H_{40}BBrN_{4}O_{5}S$: C, 54.64; H, 6.11; N, 8.50; B, 1.64.

10 Found: C, 54.52; H, 6.16; N, 8.45; B, 1.60.

K: $[a]_D = -15.07^\circ$ (c = 0.604, MeOH); ¹H NMR (300 MHz, DMSO - d₆) 9.98 (1 H, br s), 8.96 (4 H, br s), 7.93 (1 H, narrow m), 7.80 (1 H, app d), 7.64 (1 H, m), 7.56

- 15 (1 H, app t), 7.25 7.42 (5 H, m), 5.13 (2 H, s), 4.11 (1 H, dd, J = 8.3, 1.7), 3.30 (3 H, s), 3.10 -3.25 (2 H, m), 2.57 - 2.68 (1 H, m), 2.15 - 2.30 (1 H, m), 1.97 - 2.10 (1 H, m), 1.48 - 1.90 (7 H, m), 1.44 (1 H, d, J = 9.9), 1.31 (3 H, s), 1.24 (3 H, s),
- 20 0.84 (3 H, s); IR (KBr) 2500 3700 (br), 1710, 1647, 1159 cm⁻¹; MS (CI NH₃), m/e (%) 593.2 (1.2, (M + H)⁺), 568.3 (22, (M + NH₄ H₂NCN)⁺), 551.3 (100, (M + H H₂NCN)⁺); Anal. Calcd for C₃₁H₄₂BBrN₄O₅S: C, 55.29; H, 6.29; N, 8.32; B, 1.61. Found: C, 55.15;
- 25 H, 6.21; N, 8.22; B, 1.47.

L: $[a]_D = -14.12^\circ$ (c = 0.602, MeOH); ¹H NMR (300 MHz, DMSO - d₆) 10.09 (1 H, br s), 8.98 (4 H, br s), 7.90 (2 H, d, J = 8.3), 7.42 (2 H, d, J = 8.3), 4.06 (1 H, d, J = 7.0), 3.15 - 3.20 (2 H, m), 2.70 (2 H, q, J = 7.7), 2.54 (1 H, m), 2.18 - 2.28 (1 H, m), 1.98 - 2.08 (1 H, m), 1.44 - 1.84 (8 H, m), 1.30 (3 H, s), 1.24 (3 H, s), 1.20 (3 H, t, J = 7.7), 0.84 (3 H, s); IR (KBr) 2600 - 3700 (br), 1646, 1614, 1598, 1570, 35 1500, 1123 cm⁻¹; MS (DCI - NH₃), m/e (%) 458 (100, (M)

+ H) +); Anal. Calcd for C₂₄H₃₇BBrN₃O₃S: C, 53.54; H, 6.93; N, 7.81; B, 2.01. Found: C, 53.75; H, 6.98; N, 7.74; B, 1.97.

- M: [a]_D = -14.21° (c = 0.556, MeOH); ¹H NMR (300 MHz, DMSO d6) 10.06 (1 H, br s), 8.95 (4 H, br s), 7.88 (2 H, d, J = 8.1), 7.40 (2 H, d, J = 8.1), 4.06 (1 H, dd, J = 1.7, 8.3), 3.14 3.17 (2 H, m), 2.65 (2 H, t, J = 7.5), 2.50 2.60 (1 H, m), 2.18 2.28 (1 H, m), 1.98 2.08 (1 H, m), 1.45 1.84 (10 H, m), 1.30 (3 H, s), 1.24 (3 H, s), 0.89 (3 H, t, J = 7.3), 0.84 (3 H, s); IR (KBr) 2500 3700 (br), 1646, 1614, 1598, 1570, 1500, 1446, 1236, 1124, 1082 cm⁻¹; MS (CI NH₃), m/e (%) 472.2 (13.5, (M + H)⁺), 430.2 (100, (M + H H₂NCN)⁺), 278.0 (61.9); Anal. Calcd for C₂₅H₃₉BBrN₃O₃S: C, 54.36; H, 7.12; N, 7.61; B, 1.96.
- N: $[a]_D = -13.79^\circ$ (c = 0.602, MeOH); ¹H NMR (300 MHz, 20 DMSO d₆) 10.03 (1 H, br s), 8.94 (4 H, br s), 7.89 (2 H, d, J = 8.3), 7.45 (2 H, d, J = 8.3), 4.06 (1 H, br d), 3.10 3.23 (2 H, m), 2.90 3.05 (1 H, m), 2.50 2.60 (1 H, m), 2.15 2.30 (1 H, m), 1.95 2.08 (1 H, m), 1.42 1.89 (8 H, m), 1.30 (3 H, s), 1.24 (3 H, s), 1.23 (6 H, d, J = 7.0), 0.84 (3 H, s); IR (KBr) 2500 3700 (br), 1646, 1613, 1598, 1123 cm⁻¹; MS (DCI NH₃), m/e (%) 472 (100, $(M + H)^+$), 430

Found: C, 54.50; H, 7.18; N, 7.83; B, 1.73.

30 Found: C, 54.64; H, 7.17; N, 7.50; B, 1.74.

(37, (M + H - H₂NCN)⁺); Anal. Calcd for

O: $[a]_D = -13.19^\circ$ (c = 0.364, MeOH); ¹H NMR (300 MHz, DMSO - d₆) 10.03 (1 H, br s), 8.93 (4 H, br s), 7.88 (2 H, d, J = 8.5), 7.40 (2 H, d, J = 8.5), 4.06 (1 H, br d, J = 6.6), 3.15 - 3.20 (2 H, m), 2.67 (2 H, t, J

C₂₅H₃₉BBrN₃O₃S: C, 54.36; H, 7.12; N, 7.61; B, 1.96.

= 7.7), 2.50 - 2.60 (1 H, m), 2.18 - 2.28 (1 H, m), 1.95 - 2.08 (1 H, m), 1.24 - 1.84 (10 H, m), 1.23 - 1.35 (2 H, m), 1.30 (3 H, s), 1.24 (3 H, s), 0.90 (3 H, t, J = 7.3), 0.84 (3 H, s); IR (KBr) 2500 - 3700 (br), 1646, 1614, 1598, 1500, 1123 cm⁻¹; MS (CI - NH₃), m/e (%) 486.2 (3.3, (M + H)⁺), 444.2 (87.1, (M + H - H₂NCN)⁺), 292.0 (100); Anal. Calcd for C₂₆H₄₁BBrN₃O₃S: C, 55.13; H, 7.30; N, 7.42; B, 1.91. Found: C, 54.99; H, 7.22; N, 7.29; B, 2.07.

10 P: $[a]_D = -12.71^\circ$ (c = 0.598, MeOH); ¹H NMR (300 MHz, DMSO - d₆) 10.05 (1 H, br s), 8.95 (4 H, br s), 7.90 (2 H, d, J = 8.6), 7.59 (2 H, d, J = 8.6), 4.06 (1 H, br d), 3.10 - 3.23 (2 H, m), 2.50 - 2.62 (1 H, m),

15 2.16 - 2.30 (1 H, m), 1.96 - 2.08 (1 H, m), 1.42 - 1.90 (8 H, m), 1.31 (9 H, s), 1.30 (3 H, s), 1.24 (3 H, s), 0.84 (3 H, s); IR (KBr) 2500 - 3700 (br), 1646, 1613, 1597, 1498, 1123 cm⁻¹; MS (DCI - NH₃), m/e (%) 486 (100, (M + H)⁺), 444 (16, (M + H -

20 H₂NCN)⁺); Anal. Calcd for C₂₆H₄₁BBrN₃O₃S: C, 55.13; H, 7.30; N, 7.42; B, 1.91. Found: C, 55.09; H, 7.45; N, 7.40; B, 1.67.

Q: H NMR (300 MHz, DMSO - d₆) \$10.06 (1 H, br s), 8.95 (4 H, br s), 7.88 (2 H, d, J = 8.5), 7.40 (2 H, d, J = 8.5), 4.06 (1 H, br d, J = 6.6), 3.10 - 3.23 (2 H, m), 2.66 (2 H, t, J = 7.7), 2.50 - 2.60 (1 H, m), 2.15 - 2.30 (1 H, m), 1.95 - 2.08 (1 H, m), 1.40 - 1.90 (10 H, m), 1.20 - 1.38 (12 H, m), 0.80 - 0.90 (6 H, m); IR (KBr) 2500 - 3700 (br), 1646, 1614, 1598, 1500, 1124 cm⁻¹; MS (DCI - NH₃), m/e (%) 514 (100, (M + H)⁺), 472 (16, (M + H - H₂NCN)⁺); Anal. Calcd for $C_{28}H_{45}BBrN_{3}O_{3}S$: C, 56.57; H, 7.63; N, 7.07; B, 1.82. Found: C, 56.19; H, 7.53; N, 6.97; B, 1.99.

R: $[a]_D = -11.70^\circ$ (c = 0.530, MeOH); ¹H NMR (300 MHz, DMSO - d₆) d10.05 (1 H, br s), 8.83 - 9.13 (4 H, br d), 7.88 (2 H, d, J = 8.3), 7.43 (2 H, d, J = 8.3), 4.06 (1 H, br d), 3.05 - 3.25 (2 H, m), 2.45 - 2.67 (2 H, m), 2.13 - 2.30 (1 H, m), 1.94 - 2.10 (1 H, m), 1.30 - 1.90 (18 H, m), 1.30 (3 H, s), 1.24 (3 H, s), 0.84 (3 H, s); IR (KBr) 2500 - 3700 (br), 1646, 1613, 1598, 1500, 1448, 1122 cm⁻¹; MS (DCI - NH₃), m/e (%) 512 (100, (M + H)⁺), 470 (40, (M + H - H₂NCN)⁺); Anal. Calcd for C₂₈H₄₃BBrN₃O₃S: C, 56.77; H, 7.32; N, 7.09; B, 1.82. Found: C, 56.49; H, 7.38; N, 6.96; B, 1.75. S: HRMS (DCI - NH₃), Calc: 577.3019, Found: 577.3025.

15 T: $[a]_D = -8.31^\circ$ (c = 0.614, MeOH); ¹H NMR (300 MHz, DMSO - d₆) d9.98 (1 H, br s), 8.95 (4 H, br s), 7.93 (2 H, d, J = 8.8), 7.11 (2 H, d, J = 8.8), 4.00 - 4.10 (3 H, m), 3.10 - 3.23 (2 H, m), 2.50 - 2.60 (1 H, m), 2.15 - 2.30 (1 H, m), 1.95 - 2.08 (1 H, m),

- 20 1.37 1.90 (12 H, m), 1.29 (3 H, s), 1.24 (3 H, s), 0.94 (3 H, t, J = 7.4), 0.84 (3 H, s); IR (KBr) 2500 3700 (br), 1646, 1608, 1498, 1262, 1124 cm⁻¹; MS (DCI NH₃), m/e (%) 502 (100, (M + H)⁺), 460 (28, (M + H H₂NCN)⁺); Anal. Calcd for C₂₆H₄1BBrN₃O₄S: C,
- 25 53.62; H, 7.10; N, 7.21; B, 1.86. Found: C, 53.61; H, 7.09; N, 7.20; B, 1.78.
 - U: HRMS (DCI NH₃), Calc: 513.2707, Found: 513.2702.
- 30 V: HRMS (DCI NH₃), Calc: 555.3165, Found: 555.3176. W: HRMS (DCI - NH₃), Calc: 579.2812, Found: 579.2801.
 - X: HRMS (DCI NH₃), Calc: 450.2962, Found: 450.2958.
- 35 Y: HRMS (DCI NH₃), Calc: 640.3016, Found: 640.3022.

Z: $[a]_D = -8.80^\circ$ (c = 0.602, MeOH); ¹H NMR (300 MHz, DMSO - d_6) 10.03 (1 H, br s), 9.25 (1 H, br s), 8.96 (4 H, br s), 7.92 (1 H, d, J = 1.5), 7.72 (1 H, dd, J= 8.1, 1.5), 7.25 - 7.50 (6 H, m), 5.17 (2 H, s),4.08 (1 H, dd, J = 8.1, 1.5), 3.08 - 3.27 (2 H, m), 2.65 (2 H, br t), 2.50 - 2.60 (1 H, m), 2.15 - 2.30(1 H, m), 1.95 - 2.08 (1 H, m), 1.40 - 1.90 (10 H, m)m), 1.30 (3 H, s), 1.24 (3 H, s), 1.15 - 1.38 (2 H, m, buried underneath methyl absorptions), 0.77 - 0.95 10 (6 H, m); IR (KBr) 2500 - 3700 (br), 1704, 1646, 1572, 1539, 1453, 1234, 1123, 1056 cm^{-1} ; MS (CI - NH_3), m/e (%) 593.2 (1.3, $(M + H - H_2NCN)^+$), 485.2 (42.7), 333.0 (100); Anal. Calcd for C₃₄H₄₈BBrN₄O₅S: C, 57.07; H, 6.76; N, 7.83; B, 1.51. Found: C, 15

()

AA: 1 H NMR (300 MHz, DMSO - d₆) §9.98 (1 H, br s), 8.98 (4 H, br s), 7.77 - 7.92 (2 H, m), 7.08 - 7.55

57.17; H, 6.84; N, 7.76; B, 1.41.

- 20 (6 H, m), 4.90 5.30 (2 H, m), 4.09 (1 H, br d), 3.04 3.35 (5 H, m), 2.35 2.65 (3 H, m), 2.15 2.30 (1 H, m), 1.97- 2.10 (1 H, m), 1.37- 1.93 (10 H, m), 1.31 (3 H, s), 1.24 (3 H, s), 1.10 1.37 (2 H, m, buried underneath methyl absorptions), 0.72 0.93
- 25 (6 H, m); MS (CI NH₃), m/e (%) 649.4 (1.9, (M + H)⁺), 624.4 (31, (M + NH₄ H₂NCN)⁺), 607.2 (100, (M + H H₂NCN)⁺), 455.0 (39), 444.0 (29.8); Anal. Calcd for C₃₅H₅₀BBrN₄O₅S: C, 57.62; H, 6.91; N, 7.68; B, 1.48. Found:
- 30 C, 57.37; H, 6.86; N, 7.64; B, 1.40. BB: HRMS (DCI - NH₃), Calc: 520.2805, Found: 520.2796.
 - CC: HRMS (DCI NH₃), Calc: 560.2390, Found: 560.2407.
- 35 DD: HRMS (DCI NH₃), Calc: 596.2060, Found: 596.2055.

EE: HRMS (DCI - NH₃), Calc: 546.2597, Found: 546.2604.

FF: HRMS (DCI - NH₃), Calc: 534.2597, Found: 534.2609.

5 GG: HRMS (DCI - NH₃), Calc: 532.2441, Found: 532.2445.

HH: HRMS (DCI - NH₃), Calc: 532.2441, Found: 532.2452.

II: HRMS (DCI - NH₃), Calc: 480.2493, Found: 480.2492.

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JJ: HRMS (DCI - NH₃), Calc: 496.2441, Found: 496.2449.

KK: HRMS (DCI - NH₃), Calc: 507.2601, Found: 507.2592.

15 LL: HRMS (DCI - NH₃), Calc: 537.2667, Found: 537.2685.

MM: HRMS (DCI - NH₃), Calc: 498.2233, Found: 498.2231.

NN: HRMS (DCI - NH₃), Calc: 481.2445, Found: 481.2442.

20

OO: HRMS (DCI - NH₃), Calc: 557.2758, Found: 557.2754.

PP: HRMS (DCI - NH₃), Calc: 5481.2445, Found: 481.2440.

25 QQ: HRMS (NH3) - CI/DEP), Calc: 503.3193, Found: 503.3199.

RR: HRMS (DCI-NH3), Calc: 605.333; Found: 605.3325.

30 Utility

The compounds of formula (I) are useful as inhibitors of trypsin-like enzymes, notably human thrombin, plasma kallikrein and plasmin. Because of their inhibitory action, these compounds are indicated for use in the prevention or treatment of physiological

reactions catalyzed by the aforesaid enzymes such as blood coagulation and inflammation.

As an illustration of the above, the biological activity of compounds of the present invention is demonstrated by their *in vitro* inhibition of synthetic substrate hydrolysis by human thrombin S-2238 Chromogenic Assay (IC₅₀). The synthetic substrate H-D-Phe-Pip-Arg-pNA (S-2238, Kabi) is cleaved by thrombin, liberating the *p*-nitroanalide group which absorbs light at 405 nm. Enzyme activity is measured in both the presence and absence of inhibitor. A decrease in absorbance at 405 nm in the presence of inhibitor is indicative of thrombin inhibition.

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A mixture of 10 μL human thrombin (Enzyme Research Laboratories, Inc.) at an activity of approximately 7 units/mL, 10 μL of the inhibitor (normally at a concentration of 10⁻³ M or less), and 160 μL buffer (0.15 M NaCl, 10 mM HEPES, 10 mM Tris, 1 g/L PEG 8,000, pH 7.4) are incubated for 10 minutes at room temperature. To this mixture is added 20 μL of the synthetic substrate S-2238 at a concentration of 1 mM and the reaction allowed to occur for 10 minutes, after which absorbance at 405 nm is determined.

Using the methodology described above, representative compounds of this invention were evaluated and found to exhibit an IC50 of less than 1 mM, thereby confirming the utility of the compounds of the invention as effective thrombin inhibitors.

Since the compounds of formula (I) have antithrombogenic properties, they may be employed when an
anti-thrombogenic agent is indicated, such as for
control of the coagulation or the fibrinolysis system
in mammals or they may be added to blood for the
purpose of preventing coagulation or the blood due to

contact with blood collecting or distribution containers, tubing or apparatus.

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this invention.

Generally, these compounds may be administered orally or parenterally to a host to obtain an antithrombogenic effect. The dosage of the active compound depends on the mammalian species, body weight, age, and mode of administration as will be obvious to one skilled in the art. In the case of large mammals such as humans, the compounds may be administered alone or in combination with pharmaceutical carriers or diluents at a dose of from 0.02 to 15 mg/Kg to obtain the anti-thrombogenic effect, and may be given as a single dose or in divided doses or as a sustained release formulation.

Pharmaceutical carriers or diluents are well known and include sugars, starches and water, which may be used to make tablets, capsules, injectable solutions or the like which can serve as suitable dosage forms for administration of the compounds of this invention. Remington's Pharmaceutical Sciences, A. Osol, is a standard reference text which discloses suitable pharmaceutical carriers and dosage forms. The disclosure of this text is hereby incorporated by reference for a more complete teaching of suitable dosage forms for administration of the compounds of

WHAT IS CLAIMED IS:

A compound of formula (I)

 $R^{1}-Z-CCHR^{2}-BY^{1}Y^{2}$

5

(I)

wherein

 Y^1 and Y^2 are independently

- a) -OH
- 10
- b) -F,
- c) NR^3R^4 , or
- d) C1-C8- alkoxy;

 Y^1 and Y^2 when taken together can form

- a) a cyclic boron ester where said chain or ring contains from 2 to 20 carbon atoms and, optionally, a heteroatom which can be N, S, or O,
 - a divalent cyclic boro amide where said chain or ring contains from 2 t 20 carbon atoms,
- 20 c) a cyclic boro amide-ester where said chain or ring contains from 2 to 20 carbon atoms;
 - Z is
 - a) $-(CH_2)_mCONR^8-$
- 25
- b) -(CH₂)_mCSNR⁸-,
- c) -(CH₂)_mSO₂NR⁸-,
- d) $-(CH_2)_mCO_2-$,
- e) $-(CH_2)_mC(S)O-$, or
- f). $-(CH_2)_mSO_2O-;$

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 \mathbb{R}^1 is

a) -(CH₂)_p-aryl, wherein aryl is phenyl, naphthyl or biphenyl substituted with one, two or three substituents selected from the group consisting of halo (F, Cl, Br, I), CN, Cl-Cl0-alkyl, C3-C8-cycloalkyl, C2-Cl0-alkenyl,

C2--C10-alkynyl, $-R^8$, $-OR^8$, methylenedioxy, $-NO_2$, $-CF_3$, $-S(O)_rR^7$, NR^8R^9 , $-CO_2R^8$, $-CO_2R^$

- b) heteroaryl, wherein heteroaryl is an unsubstituted or monosubstituted or disubstituted
 - i) 5- or 6-membered aromatic ring, which contains from 1 to 3 heteroatoms selected from the group consisting of O, N, and S,
 - ii) quinolinyl,
 - iii) isoquinolinyl,
 - iv) benzopyranyl,
 - v) benzothiophenyl,
- 15 vi) benzofuranyl,

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- vii) 5,6,7,8-tetrahydroquinolinyl
- viii) 5,6,7,8-tetrahydroisoquinolinyl

and wherein the subtitutents are members selected from the group consisting of halo (F, Cl, Br, I, -CN, Cl-Cl0-alkyl, C3-C8-cycloalkyl, C2-Cl0-alkenyl, C2-Cl0-alkynyl, -R⁸, OR⁸, NO₂, -CF₃, -S(O)_rR⁷, NR⁸R⁹, -COR⁸, -CONR⁸R⁹, NR⁸COR⁹, NR⁸CO2R⁹,

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d) R¹⁰

e) .\{\begin{align*}
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•

f)

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ξ (CH₂)_t

10 g)

2 O RIV

 \mathbb{R}^2 is

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a) $-(CH_2)_n$ -NHC (NH) NH₂,

b) $-(CH_2)_n$ -NHC (NH) NHCOCH3,

c) $-(CH_2)_n-SC(NH)NH_2$,

e) $-(CH_2)_n-SC(NH)_2$, or

f) - (CH) n-NH (2-pyridyl);

R³ is H, phenyl or C1-C4-alkyl;

 ${\ensuremath{\mathbb{R}}}^4$ is H, or phenylsulfonyl;

- ${\rm R}^5$ and ${\rm R}^6$ are hydrogen or when taken together form a six membered aromatic ring optionally substituted with one, two or three substituents selected from the
- group consisting of halo (F, C1, Br, I), -CN, C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl, C2-C10-alkynyl, -OR⁸, -NO₂, -CF₃, -S(O)_rR⁷, -NR⁸R⁹, -COR⁸, -COR₂R⁸, -CONR⁸R⁹, phenyl, benzyl, phenylethyl;

 R^7 is

- 10 a) phenyl,
 - b) C1-C4-alkyl,
 - c) C1-C4-alkoxy, or
 - d) -CF3;

 \mathbb{R}^8 and \mathbb{R}^9 are independently

- 15 a) H,
 - b)

- c) C3-C7-Cycloalkyl,
- d) C1-C8-alkyl;
- 20 R^{10} and R^{11} are independently
 - a) halo (F, C1, Br, I),
 - b) -CN,
 - c) C1-C10-alkyl,
 - d) C3-C8-cycloalkyl,
- e) C2-C10-alkenyl,
 - f) C2-C10-alkynyl,
 - $q) OR^8$
 - h) NO2,
 - i) -CF3,
- 30 j) $-s(0)_{r}R^{7}$,
 - $k) -NR^8R^9$
 - 1) $-COR^9$,
 - m) $-CO_2R^8$, or
 - n) $-CONR^8R^9$;

 R^{12} is

- a) H,
- b) C1-C4-alkyl,
- 5 c) phenyl
 - d) benzyl,
 - e) - COR^7
 - $f) -SO_2R^7$

m is 0 to 6;

- 10 n is 3 or 4;
 - p is 0 to 2;
 - r is 0 to 2;
 - t is 1 to 5

E is -CO-, $-SO_2-$, $-CH_2-$ or a single bond,

15 F is -CO-, and pharmaceutically acceptable salts thereof.

2. A compound of Claim 1 wherein:

R1 is phenyl containing 1-3

substituents selected from the series halo (F, CL, Br, I), C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl, C2-C10-alkynyl, $-R^8$, $-OR^8$, $-NO_2$, $-CF_3$, $-S(O)_rR^7$, $-NR^8R^9$, $-COR^8$, $-CO_2R^8$, $CONR^8R^9$, NR^8COR^9 , and

-ξ-__NR¹²

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(;

R₂ is

- a) -(CH₂)₃-NHC(NH)NH₂, or
- b) $-(CH_2)_3-SC(NH)_NH_2$.
- 30 3. A compound of Claim 2 wherein Z is $-(CH_2)_mCONR^8-$.
 - 4. A compound of Claim 3 selected from the group consisting of

 N^{1} -(4-phenylbenzoyl)-(R)-boroarganine, hydrochloride,

 N^{1} -(3-phenoxybenzoyl)-(R)-boroarganine, hydrochloride,

- N^{1} -(1-fluorenonyl)-(R)-boroarginine, hydrochloride,
- N^{1} -(4-[buty1]benzoy1)-(R)-boroarginine, hydrochloride,
- N¹-(2-benzoylbenzoyl)-R-boroarginine, hydrochloride,
- N¹-(5-phenyl-2-furol)-R-boroarginine, hydrochloride,
 - N¹-(3-[N-benzyloxycarbonyl-N-methylamino]-4-[1-butyl]-benzoyl)-(R)-boroarginine, hydrochloride,
 - N¹-(2-phenyl-4-isoquinolyl)-(R)-boroarginine, hydrochloride,
- 10 N¹-(4-cyclohexylbenzoyl)-(R)-boroarginine, hydrochloride
 - N1-(2-methyl-4-phenylbenzoyl)-(R)-boroarginine, hydrochloride, or
- 5. A pharmaceutical composition comprising a

 pharmaceutically suitable carrier and a
 therapeutically effective amount of a compound of any
 one of claims 1 through 4.
- 6. A method of treating a physiological disorder in a
 warm blooded animal catalyzed by trypsin-like enzymes
 comprising administering to an animal in need of such
 treatment an effective amount of a compound of any
 one of claims 1 through 4.

INTERNATIONAL SEARCH REPORT

Inte mal Application No PCT/US 94/02965

| A. CLASS | IFICATION OF SUBJECT MATTER C07F5/02 A61K31/69 | | |
|--|--|--|--|
| According to | to International Patent Classification (IPC) or to both national class | ification and IPC | |
| | SEARCHED | | |
| | locumentation searched (classification system followed by classifica- | tion symbols) | |
| IPC 5 | CO7F A61K | | , |
| Documentat | tion searched other than minimum documentation to the extent that | such documents are included in the fields a | searched |
| Electronic d | ata hase consulted during the international search (name of data ba | se and, where practical, search terms used) | |
| C. DOCUM | HENTS CONSIDERED TO BE RELEVANT | | |
| Category * | Citation of document, with indication, where appropriate, of the r | elevant passages | Relevant to claim No. |
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| A | WO,A,92 07869 (KAKKAR, V.V. ET Al 1992 see the whole document | L.) 14 May | 1-6 |
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| Furt | ner documents are listed in the continuation of box C. | X Patent family members are listed | in annex. |
| "A" docume consider filing docume which in citation "O" docume other n" "P" docume later the | ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or | T later document published after the into or priority date and not in conflict with cited to understand the principle or the invention "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the default of particular relevance; the cannot be considered to involve an involve an inventity is combined with one or ments, such combined with one or ments, such combination being obvious the art. "&" document member of the same patent. Date of mailing of the international se | th the application but hecory underlying the claimed invention to be considered to between is taken alone daimed invention the top when the top when the top when the top other such docuurs to a person stolled |
| | June 1994 | 14.06.9 | |
| Name and m | nailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 FIV Rijswijk Tel. (+ 31-70) 340-2040, Tx. 31 651 cpo nl, Fax (+ 31-70) 340-3016 | Authorized officer Rinkel, L | |

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US 94/02965

| This in | ternational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons: |
|----------|---|
| 1. | Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: "Remark: Although claim 6 is directed to a method of treatment of |
| _ | (diagnostic method practised on) the human/animal body the search has been carried out and based on the alleged effects of the compound/composition." |
| 2 | Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: |
| | |
| 3. [| Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). |
| Box II | Observations where unity of invention is lacking (Continuation of item 2 of first sheet) |
| This Int | ernational Searching Authority found multiple inventions in this international application, as follows: |
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| | |
| . 🗆 | As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. |
| . 🗌 | As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee. |
| | |
| · 📋 | As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: |
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| | No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: |
| | No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: |
| | No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: The additional search fees were accompanied by the applicant's protest. |

INTERNATIONAL SEARCH REPORT

information on patent family members

Int onal Application No PCT/US 94/02965

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